The role of prenatal and perinatal factors in eating disorders:
a systematic review

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Abstract
Numerous studies showed that factors influencing fetal development and neonatal period could lead to lasting alterations in the brain of the offspring, in turn increasing the risk for eating disorders (EDs). This work aims to systematically and critically review the literature on the association of prenatal and perinatal factors with the onset of EDs in the offspring, updating previous findings and focusing on anorexia nervosa (AN) and bulimia nervosa (BN). A systematic literature search was performed on Pubmed, PsycINFO, and Scopus. The drafting of this systematic review was conducted following the PRISMA statement criteria and the methodological quality of each study was assessed by the MMAT 2018. A total of 37 studies were included in this review. The factors that showed a more robust association with AN were higher maternal age, preeclampsia and eclampsia, multiparity, hypoxic complications, prematurity, or being born preterm (< 32 weeks) and small for gestational age or lower birth size. BN was only associated with maternal stress during pregnancy. Many methodological flaws emerged in the considered studies, so further research is needed to clarify these inconsistencies. Altogether, data are suggestive of an association between prenatal and perinatal factors and the onset of EDs in the offspring. Nevertheless, given the methodological quality of the available literature, firm conclusions cannot be drawn and whether this vulnerability is specific to EDs or mental disorders remains to be defined. Also, a strong need for longitudinal and well-designed studies on this topic emerged.

Keywords Eating disorders · Anorexia nervosa · Bulimia nervosa · Pregnancy complications · Obstetric complications

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Introduction
Eating disorders (EDs) are complex mental illnesses characterized by unknown etiology, with many putative risk factors (Fairburn and Harrison 2003; Dalle Grave 2011; Jacob et al. 2011) facilitating the ED onset. Genetic risk factors are of great importance, with genes impacting on the development of both anorexia nervosa (AN) and bulimia nervosa (BN) as well as on their predisposing traits (Trace et al. 2013; Baker et al. 2017). An altered neurodevelopment has been implicated in the pathogenesis of several mental disorders (Katzman et al. 1997; Chowdhury et al. 2003; King et al. 2018) such as schizophrenia (Geddes and Lawrie 1995; Verdoux et al. 1997; Geddes et al. 1999; Cannon et al. 2000), attention deficit hyperactivity disorder (Lindström et al. 2006), and autism (Gardener et al. 2009). With more detail, hypoxic complications and prematurity have been associated with schizophrenia risk (Geddes and Lawrie 1995; Verdoux et al. 1997; Geddes et al. 1999; Cannon et al. 2000), leading to the formulation of a “neurodevelopmental hypothesis for schizophrenia” (Rapoport et al. 2012).
Several studies (Gillberg et al. 1994; Connan et al. 2003) linked prenatal and perinatal complications (Raevuori et al. 2014) to EDs, with the hypothesis that subtle neurological damages (Gillberg et al. 1994), neuropsychological disabilities (Galdersi et al. 2003), and nonreversible morphological brain changes could ease the disorder onset. Krug and co-workers (Krug et al. 2013) systematically reviewed the literature on obstetric complications (OCs) and EDs, selecting 14 articles, and performing a meta-analysis where possible. Conflicting results emerged, so their meta-analysis found a nonsignificant association between instrumental delivery and prematurity and ED risk. More recently, a descriptive review of 22 articles focusing on additional risk factors (e.g., the role of sex hormones, maternal status, and maternal EDs) found once more mixed results (Raevuori et al. 2014). Another review of 13 studies (Jones et al. 2017) focused instead on “fetal programming” as a model on how stimuli/insults occurring during critical or sensitive periods of fetal development could have physiological effects that unfold across the life span. The latter review introduced new risk factors, such as maternal stress during pregnancy, but finding again controversial results. Therefore, the aim of this systematic review is twofold: (a) to critically review the updated literature on the association of prenatal and perinatal factors with the onset of EDs in the offspring and (b) to expand knowledge on the critical points that need to be addressed by future lines of research.

Methods

Search strategy and selection criteria

The drafting of this systematic review was conducted following the PRISMA statement criteria (Moher et al. 2009), and the studies included in this systematic review have been evaluated by the MMAT (Mixed Methods Appraisal Tool, 2018 version; Hong et al. 2018).

A systematic literature search was done between March 1 and May 1, 2019, including online database searches, namely, Pubmed, PsycINFO, and Scopus, and journal hand searching to ensure a wide inclusion of eligible studies.

The search was designed to include those studies published since 1998; this is a reasoned choice as the purpose of this systematic review was to analyze the most recent studies on the topic. All full-text studies published were included if they met the following inclusion criteria:

- Study type criteria: cohort study and case-control study
- Eating disorder assessment criteria (EDs): DSM (III, III-R, IV, IV-TR, 5) diagnosis, ICD (8,9,10) diagnosis, clinical diagnosis or self-report diagnosis, or ED symptomatology obtained with questionnaire scores
- Risk assessment criteria: obstetric birth records, parents recall, and/or a combination of both

The following list of terms were included and combined together in different search lines: “eating disorders,” “anorexia nervosa,” “bulimia nervosa,” “pregnancy complications,” “obstetric complications,” “prenatal risk factors,” “perinatal risk factors,” “gestational diabetes,” “hypertensive disease in pregnancy,” “maternal anemia,” “preeclampsia,” “ eclampsia,” “maternal viral infection,” “season of birth,” “vit. D levels during pregnancy,” “prenatal sex hormones,” “dysmaturity,” “preterm birth,” “mother weight,” “mother smoke habit,” “maternal stress,” “maternal anxiety,” “placenta previa,” “pregnancy bleeding,” “breech delivery,” “induced labor,” “inertia uteri,” “premature rupture of the membrane,” “forceps,” “cesarean section,” “vaginal instrumental delivery,” “vacuum extraction,” “cephalohematoma,” “umbilical cord wrapped around the neck,” “placental infarction,” “cyanosis,” “jaundice,” “need for resuscitation,” “need for oxygen,” “need for intubation,” “birth weight,” “prematurity,” “tremors,” “hypothermia,” “hypotonia,” and “neuromuscular disturbances.”

Two investigators (E.M., F.C.) screened the title and abstracts of 851 studies identified through the search using the inclusion criteria. Full-text articles were retrieved for all studies that met the inclusion criteria or required more information than was provided in the abstract. The reasons for excluding the studies were documented and shown in Fig. 1.

Data analysis

All database search results were imported into Zotero. Duplicate entries were removed before the screening. Data from the studies were extracted and summarized (see Table 1 and Supplementary Materials). Data extracted included authors, year, study type, sample size, psychiatric assessment, maternal factors, pregnancy complications, obstetric complications, neonatal factors, and main findings.

As reported in Table 1, the methodological quality of each study was assessed by two investigators (F.C. and E.M.) using MMAT (Mixed Methods Appraisal Tool, 2018 version; Hong et al. 2018) that allows a quality assessment of qualitative, quantitative, and mixed methods studies (Pluye et al. 2009; Pace et al. 2012) with a double evaluation and a high intraclass correlation. The overall agreement was 90% on the MMAT items; when discrepancies emerged, they were resolved through discussion. The quality score ranged from 0 (no criteria met) to 5 (all criteria met).

According to earlier literature (Whittemore and Knafl 2005), due to the extreme heterogeneity of the study results, it has not been possible to conduct a meta-analysis so it was preferred to proceed with a thematic meta-synthetic approach to critically synthesize literature on the role of prenatal and
perinatal factors on the onset of EDs. The findings of all the included studies were independently read and re-read, coded, and organized into categories, which were then compared across studies to identify relationships and themes (Whittemore and Knafl 2005).

**Results**

Of the 815 studies screened, 96 full-text articles were assessed for eligibility, and 37 were finally included in the systematic review (please see Table 1 and Supplementary Materials for all details).

The outcomes assessed in the studies were highly variable and were divided as follows: (1) maternal factors, (2) pregnancy complications, (3) obstetric complications, and (4) neonatal factors. Study distribution has been outlined in Fig. 2.

As shown in Fig. 3, the methodological quality of studies ranged from low (2 points) to high (5 points). Overall, no studies were rated as showing very low methodological quality (1 point) and only the category of pregnancy complications showed the majority of studies (56%) as scoring low and fair. In fact, all other categories (pregnancy complications, obstetric complications, and neonatal factors) reported about 60% of studies with high (4 points) or very high (5 points) methodological quality scores (see Fig. 3).

The most frequent limitations were participants’ poor representativeness of the target population (Shoebridge and Gowers 2000; Foley et al. 2001; Feingold et al. 2002; Klump et al. 2006; Culbert et al. 2008, 2013; Smith et al. 2010; Coombs et al. 2011; Quinton et al. 2011; Nosarti et al. 2012; St-Hilaire et al. 2015), small sample size (Shoebridge and Gowers 2000; Feingold et al. 2002; Klump et al. 2006; Culbert et al. 2008, 2013; Smith et al. 2010; Quinton et al. 2011), and unappropriate measurements (e.g., self-report diagnosis, ED symptomatology investigated with questionnaires on healthy participants, lack of a clear DSM or ICD diagnosis; Feingold et al. 2002; Montgomery et al. 2005; Klump et al. 2006; Culbert et al. 2008, 2013, 2015; Baker et al. 2009; Nicholls and Viner 2009; Smith et al. 2010; Wehkampli et al. 2010; Coombs et al. 2011; Quinton et al. 2011; Lydecker et al. 2012; Nosarti et al. 2012; Allen et al. 2013; Romero-Martinez and Moya-Albiol 2014; St-Hilaire et al. 2015; Lofrano-Prado et al. 2015; Matinolli et al. 2016; Sacks et al. 2016), and uncontrolled confounders (Shoebridge and Gowers 2000; Foley et al. 2001; Feingold et al. 2002;
| Authors                  | Study type          | Sample                                      | Maternal factors | Pregnancy complications | Obstetric complications                                                                 | Neonatal factors                                                                 | Main findings                                                                                      | Quality rating |
|-------------------------|---------------------|---------------------------------------------|------------------|-------------------------|-----------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|---------------|
| Cnattingius et al. 1999 | Case-control study  | AN (n = 781), HCs (n = 3905)                | Maternal age     | Hypertensive disease,   | Inertia uteri, forceps, or vacuum delivery, preterm rupture of the membranes, cesarean section, cephalohematoma, other trauma | Gestational age, birth weight and birth weight for gestational age, Apgar score, jaundice   | AN diagnosis was associated with maternal age, very preterm birth (< or = 32 weeks), small for gestational age in very preterm birth, cephalohematoma | 5             |
| Morgan et al. 2000      | Case-control study  | BN (n = 935), patients with AN history and BN diagnosis (n = 227) general population data | –                | –                       | –                                                                                       | –                                                                                           | BN diagnosis is not associated with a specific month of birth AN history and BN diagnosis are associated with a peak season of birth in March | 4             |
| Shoebridge & Gowers et al. 2000 | Case-control study | AN (n = 40) and HCs (n = 40) | –                | Antepartum hemorrhage  | Toxemia, previous obstetric complications, forceps delivery, cesarean section | –                                                                                           | No association was found | 2             |
| Foley et al. 2001       | Twin cohort study   | AN (n = 7), broadly defined AN (bdAN) (n = 71), BN (n = 42), broadly defined BN (bdBN) (n = 100), twin control (n = 158/6) | –                | High blood pressure,   | Premature contractions, labor lasting more than 24 h, breech delivery, cesarean delivery, forceps delivery, cord wrapped around the neck, blue at birth, required an incubator, any perinatal complication | Birth weight, gestational age, jaundice, failure to breathe at first, convulsions, blood transfusion | AN: low gestational age, # prematernal maternal complications bdAN: prematernal maternal complications BN: # prematernal maternal complications bdBN: # prematernal maternal complications | 3             |
| Feingold et al. 2002    | Historical prospective cohort (Thomas Jefferson University Hospital 1979–1981, n = 86) | Infants born preterm (n = 84) | Mother older than 40 years, alcohol abuse | Preeclampsia, eclampsia, chorioamnionitis, urinary tract infection, cervicitis, asthma, hyperemesis, gravidarum, cervical cancer, vaginal bleeding, multiparity | Premature rupture of the membranes, cesarean delivery, breech or transverse delivery, abruptio placenta, placenta previa, incompetent cervix, the total number of complications | Small for gestational age | No association with ED symptomatology for any of the studied variables | 4             |
| Lindberg and Hjern 2003 | Population cohort study (Swedish Inpatient Register—1973–1984) | AN (n = 1122), HCs (n = 988,749) | Maternal age | Preeclampsia | Premature rupture of the membranes, placental | Gestational age, size for gestational age, AN diagnosis was associated with maternal age | AN diagnosis was associated with maternal age | 4             |
| Authors                  | Study type               | Sample                                      | Maternal factors                                                                 | Pregnancy complications                                                                 | Obstetric complications                                    | Neonatal factors                                                                 | Main findings                                                                                   | Quality rating |
|-------------------------|--------------------------|---------------------------------------------|-----------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------------------------------------------|---------------------------------------------------------------------------------------------|---------------------------------------------------------------|----------------|
| Register Data 1973–1982 | Cohort study 1970–1982   | n = 989,871)                                | Maternal age at delivery, smoke habit, occupation, social class, psychiatric morbidity, BMI | abruption, breech delivery, cephalohematoma, neonatal O2 necessity                       | prematurity, Apgar score, distress                          | (> 25 years), gestational age at birth (23–32 weeks), preeclampsia, premature rupture of the membranes, low Apgar, O2 necessity, and distress, cephalohematoma, breech delivery | 3                                                             |
| Montgomery et al. 2005  | Cohort study 1970–1982   | BN (n = 100), HCs (3946)                    | Maternal age at delivery, smoke habit, occupation, social class                    | –                                                                                       | –                                                           | Self-reported diagnosis of BN is associated with maternal smoke during pregnancy           | 3                                                             |
| Favaro et al. 2006      | Birth cohort study 1971–1979 | AN (n = 114), BN (n = 73), HCs (n = 554)  | Maternal age, social class                                                         | Bleeding, preeclampsia, maternal diabetes, threatened miscarriages, and anemia         | Inertia uteri, breech delivery, premature rupture of the membranes, breech delivery, placental infarction or abortion, placenta previa, meconium staining of the amniotic fluid, forceps or vacuum extraction, the umbilical cord wrapped around the infant’s neck, cephalopelvic need for resuscitation O2, and intubation, disproportion, # of obstetric complications, cephalohematoma | Small for gestation age, birth weight, cyanosis, respiratory and cardiac problems, jaundice, neuromuscular disturbances such as hyporeactivity, hypotonia, and tremors, hyperthermia | 5                                                             |
| Klump et al. 2006        | Twin cohort study        | FSS (n = 113)                               | –                                                                                 | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio)                    | –                                                           | ED symptomatology was associated with lower levels of prenatal testosterone (higher 2D:4D ratio) | 2                                                             |
| Culbert et al. 2008      | Twin cohort study        | FSS (n = 304), FOS (n = 59), MOS (n = 54), MSS (n = 165), HCs (n = 69) | –                                                                                 | Levels of prenatal testosterone (indirectly studied by OS and SS twin study)          | –                                                           | Highest levels of disordered eating were observed for fSS twins, followed by fOS twins, mOS twins, and mSS twins | 3                                                             |
| Authors             | Study type                                      | Sample                                                                 | Maternal factors                         | Pregnancy complications                                                                 | Obstetric complications                                                                 | Neonatal factors                                                                 | Main findings                                                                 | Quality rating |
|---------------------|------------------------------------------------|------------------------------------------------------------------------|------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|-----------------------------------------------------------------------------|----------------|
| Favaro et al. 2008  | Birth cohort study (Padua birth cohort 1971–1979) | AN (n = 66), BN (n = 44), HCs (n = 257)                                | Maternal age                              | Vaginal bleeding, pre eclampsia, maternal diabetes, threatened miscarriages, and anemia | inertia uteri, breech delivery, premature rupture of the membranes, placental infarction or abruption, placenta previa, meconium staining of the amniotic fluid, forceps or vacuum extraction, the umbilical cord wrapped around the infant's neck, cephalohematoma, need for resuscitation | O₂ and intubation                                                            | Small for gestational age, birth weight, cyanosis, respiratory and cardiac problems, jaundice, neuromuscular disturbances such as hy poreactivity, hypotonia, and tremors, hypothermia | Preterm birth and neonatal dysmaturity were associated with high harm avoidance in the offspring affected by EDs | 4              |
| Raevuori et al. 2008| Twin cohort study (FinnTwin 16, 1975–1979 sample n = 2426) | 2426 female twins with known zygosity (OS dizygotic n = 793, SS dizygotic n = 765, monozygotic n = 868), 1,962 male twins (OS dizygotic n = 717, SS dizygotic n = 705, monozygotic n = 540) | –                                        | Levels of prenatal testosterone (indirectly studied by OS and SS twin study)            | –                                                                                       | –                                                              | Opposite-sex twin pairs were not significantly different from monozygotic or same-sex dizygotic twins (female) in the association with AN or BN | 4              |
| Baker et al. 2009   | Swedish twin study of child and adolescent development (TCHAD 1985–1986) | n = 439 identical females, n = 213 fraternal females, n = 461 identical males, n = 344 fraternal males, n = 371 opposite-gender twin pairs | –                                        | Levels of prenatal testosterone (indirectly studied by twin study)                      | –                                                                                       | –                                                              | ED symptomatology was not associated with twin types | 3              |
| Nicholls and Viner 2009 | Prospective birth cohort (British cohort study, n = 16,567) | AN (n = 101), HCs (n = 11,261)                                             | Smoke habit, ethnicity, maternal anemia     | Gestational diabetes, Perinatal hypoxia                                                   | Birth weight, gestational age, prematurity (< 37 weeks)                               | No significant differences between AN and HCs                                   |                                                                             | 3              |
| Smith et al. 2010   | Cross-sectional observational study              | Male college students (n = 204)                                           | –                                        | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio)                       | –                                                                                       | –                                                              | Higher testosterone exposure (lower 2D:4D ratio) was associated with less ED symptoms, greater drive for muscularity, diminished drive for leanness | 2              |
| Wehkalampi et al. 2010 | Case-control study (Helsinki Study of Very Low Birth Weight Adults) | Very low birth weight adults (n = 255), term (n = 189)                        | Maternal age, smoke habit, BMI, education | –                                                                                       | –                                                                                       | Gestational age, very preterm birth (< 32 weeks), very low birth weight—VLBW (< 1500 g) | In both sexes, EDI-2 scores were lower in VLBW individuals than in HCs | 3              |
| Authors          | Study type                        | Sample                                                                 | Maternal factors                                      | Pregnancy complications                     | Obstetric complications | Neonatal factors | Main findings                                                                                       | Quality rating |
|------------------|-----------------------------------|------------------------------------------------------------------------|-------------------------------------------------------|---------------------------------------------|-------------------------|------------------|----------------------------------------------------------------------------------------------------|----------------|
| Coombs et al.    | Observational study on a Midlands High School (UK) students | n = 132 pupils (age 11–14)                                              | –                                                     | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio) | –                       | –                | No strong association was found                                                                 | 2              |
| Favaro et al.    | Birth cohort study (Padua Birth Cohort 1970–1984, n = 27,682) | AN (n = 402), HCs (n = 26,950)                                          | Maternal education, maternal, and social class        | Maternal exposure during pregnancy to chickenpox, measles, rubella, or influenza | –                       | –                | Month of birth AN diagnosis was associated with the exposure to rubella and chickenpox at the 6th of pregnancy. Being born in June was associated with AN AN was associated with low 2D:4D ratio (higher prenatal testosterone) BN: high 2D:4D ratio was associated with lower prenatal testosterone for girls | 4              |
| Quinton et al.   | Case-control study                | AN (n = 25), BN (n = 26), HCs (n = 99)                                  | –                                                     | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio) | –                       | –                | Gestational age, birth weight for gestational age, newborn sex, Apgar score at 5 min | 2              |
| Nosarti et al.   | Historical population-based cohort study (Swedish Birth Register 1973–1985 and Hospital Discharge Register n = 1,301,522) | ED (n = 997), other psychiatric disorder and HCs                        | Maternal age, maternal education, maternal psychiatric family history | Parity                        | –                       | –                | Gestational age, birth weight for gestational age, newborn sex, Apgar score at 5 min | 4              |
| Lydecker et al.  | Three twin cohort study (MATR (US), NIPHTP (Norway), STAGE (Sweden)) | US: OS (n = 481), SS (n = 1,022), Norway: OS (n = 345), SS (n = 1430), Sweden: OS (n = 2,343), SS (n = 7000) | –                                                     | Levels of prenatal testosterone (indirectly studied by OS and SS twin study) | –                       | –                | No association between co-twin sex and EDs                                                                 | 4              |
| Allen et al.     | Population cohort (Western Australian Pregnancy Cohort (Raine)) n = 2900 | ED (n = 98), HCs (n = 428)                                              | Maternal age, BMI, drinking alcohol, smoking cigarettes, education, family income | Serum 25(OH)D level at 18 weeks, kidney disease or dysfunction, urinary tract infection, thyroid dysfunction | –                       | –                | Gestational age at birth, season of birth, weight, preterm birth (<37 weeks), newborn sex | 3              |

The role of prenatal and perinatal factors in eating disorders: a systematic review
| Authors                  | Study type                      | Sample                                                                 | Maternal factors                                                                 | Pregnancy complications | Obstetric complications | Neonatal factors | Main findings                                                                 | Quality rating |
|-------------------------|--------------------------------|------------------------------------------------------------------------|----------------------------------------------------------------------------------|-------------------------|--------------------------|------------------|---------------------------------------------------------------------------------|----------------|
| Culbert et al. 2013     | Twin cohort study (Michigan State Twin Study) | FSS, fOS, mOS, mSS \((n = 394),\) HCs \((n = 63)\) | –                                                                                 | Levels of prenatal testosterone (indirectly studied by OS and SS twin study) | –                         | –                | No differences were observed in the levels of disordered eating attitudes in opposite-sex and same-sex twins in pre-early puberty. During later phases of puberty, females from opposite-sex twin pairs exhibited lower disordered eating attitudes than females from same-sex twin pairs | 3              |
| Taborelli et al. 2013   | Case-control study (Sister Pair Study) | AN \((n = 94),\) BN \((n = 63),\) HCs \((n = 157)\) | Anxiety during pregnancy (questionnaire)                                        | –                       | –                         | –                | Anxiety during pregnancy was associated with AN, but not BN                   | 4              |
| Vellisa et al. 2013     | Case-control study             | AN \((n = 210),\) general population data | –                                                                                 | –                       | –                         | –                | Month of birth was not associated with the month of birth                     | 4              |
| Winje et al. 2013       | Case-control study             | AN \((n = 4045),\) HCs with the same year of birth, sex, and region of birth | –                                                                                 | –                       | –                         | –                | No significant differences between AN and HCs                                  | 4              |
| Goodman et al. 2014     | Population cohort study (Swedish Register Data, 1975–1998, \(n = 2,135,279\)) | AN \((n = 7351),\) BN \((n = 2,804),\) other eating disorders \((n = 10,408),\) HCs \((n = 2015,862)\) | Maternal age, maternal education, number of full siblings, number of half-siblings, eating disorder in mother, multiple smoke habits | Multiparity              | Premature rupture of the membranes, cesarean section, instrumental delivery, cephalohematoma, other birth trauma | Gestational age, birth weight for gestational age, birth length for gestational age, APGAR score | 5              |
| Romero-Martinez and Moya-Albiol 2014 | Observational study            | AN \((n = 34),\) HCs \((n = 40)\) | Maternal age, maternal BMI, right 2D:4D ratio, education, marital status of children | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio) | –                         | –                | Low 2D:4D ratio (higher prenatal testosterone) in AN. Salivary testosterone was negatively related to the 2D:4D ratio | 3              |
| Authors                     | Study type                  | Sample                                                                 | Maternal factors                                                                 | Pregnancy complications | Obstetric complications | Neonatal factors                                   | Main findings                                                                                                                                                                                                 | Quality rating |
|-----------------------------|-----------------------------|------------------------------------------------------------------------|----------------------------------------------------------------------------------|-------------------------|--------------------------|----------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------|
| Culbert et al. 2015         | Michigan State University Twin Registry | Study 1 (2D:4D ratios): monzygotic (n = 229) and dyzygotic (n = 180) Study 2 (OS-F study): 1538 males and females and 131 non-twin females as an additional comparison group | –                                                                                 | Levels of prenatal testosterone (indirectly studied by 2D:4D ratio and by OS-F twin study) | –                                                      | –                                                                  | In both studies, higher prenatal testosterone exposure (lower 2D:4D, females from opposite-sex twin pairs vs controls) predicted lower disordered eating symptoms in early adolescence and young adulthood | 3             |
| Lofrano-Prado et al. 2015   | Cross-sectional observational study | College students (n = 408)                                             | Mother’s age > 25 years                                                           | Number of obstetric complications, cesarean delivery | Low birth weight (<2500 g), no breastfeeding, not first in the birth order | –                                                                  | Mother’s age lower than 25 years old is associated with AN symptoms BN symptoms are associated with # of obstetric complications | 4             |
| Micali et al. 2015          | Cohort study (very preterm cohort n = 476) | VPT, < 33 weeks (n = 143)                                             | –                                                                                 | –                                                      | Cesarean section, vaginal delivery                  | Birthweight, gestational age, ventricular dilatation, neonatal complications | ED symptomatology was not associated with any of the factors. Cesarean delivery was associated with compensatory behaviors. Higher EAT-26 scores are associated with maternal exposure to stress in the third trimester | 2             |
| St-Hilaire et al. 2015      | Cohort study (Project Ice Storm cohort, n = 54) | Teenagers born from cohort mothers (n = 54)                          | Maternal age, maternal education, social class, maternal stress exposure, trimester of stress exposure (Storm 24, IES-R scales) | –                                                      | –                                                      | Birth weight, length of gestation                  |                                                                                                                                                                                                           | 3             |
| Tenconi et al. 2015         | Birth cohort study (Padua new Birth Cohort 1969–1997 and previous 1971–1979) | New cohort: AN (n = 150), BN (n = 35), HCs (n = 73) Whole cohort: AN (n = 264), BN (n = 108), HCs (n = 624) | Maternal age, social class                                                        | Bleeding, preeclampsia, maternal diabetes, threatened miscarriages, and anemia | Inertia uteri, breech delivery, premature rupture of the membranes, breech delivery, placental infarction or abruptio, placenta previa, meconium staining of the amniotic fluid, forceps or vacuum extraction, umbilical cord wrapped around the infant’s neck, and cephalopelvic disproportion, # of obstetric complications, | Small for gestation age, birth weight, cyanosis, respiratory and cardiac problems, jaundice, neuromuscular disturbances such as hyporeactivity, hypotonia, and tremors, hypothermia | AN maintains its associations as shown in Favaro (2006), and higher maternal age and weight gain during pregnancy. Maternal diabetes and anemia lost their association BN is associated with neonatal hyporeactivity, early feeding difficulties, short and small for gestational age | 5             |
| Authors                  | Study type                  | Sample                                                                 | Maternal factors                                                                 | Pregnancy complications                                                                 | Obstetric complications                                                                 | Neonatal factors                                      | Main findings                                                                 | Quality rating |
|-------------------------|-----------------------------|------------------------------------------------------------------------|----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|------------------------------------------------|-----------------------------------------------------------------------------|----------------|
| Matinolli et al. 2016   | Population cohort study     | Early preterm (n = 185), late preterm (n = 348), term-born control (n = 637) | Maternal weight, maternal smoke habit, socioeconomic status, education           | Gestational diabetes, hypertension, preeclampsia, eclampsia                                | –                                                                                        | Early preterm birth (<32 weeks) and late preterm birth (32–37 weeks)       | EDI-2 scores were significantly lower in early-born preterms than in HCs, in particular in “Body dissatisfaction” and “Drive for thinness” subscales | 4              |
| Sacks et al. 2016       | Population cohort study     | EDs (n = 486)                                                         | Maternal age, maternal obesity                                                   | Gestational diabetes mellitus (type A1 and type A2)                                      | –                                                                                        | Gestational age at birth                                                 | ED and other psychiatric disorders in the offspring were associated with gestational diabetes mellitus | 2              |
| Su et al. 2016          | Population-based cohort     | AN (n = 5878), BN (n = 1722), mixed ED (n = 3159), HCs (n = 2110,755)  | Maternal loss of a close relative 1 year prior to or during pregnancy            | –                                                                                        | –                                                                                        | –                                                                        | ED, BN, and mixed ED were associated with maternal exposure to prenatal loss | 4              |
| Razaz et al. 2018       | Retrospective Swedish cohort study | AN (n = 2414)                                                         | Maternal BMI, years of education, maternal age at delivery, smoke habit          | Vaginal delivery, vaginal instrumental delivery, elective cesarean section, emergency cesarean section | Birthweight for gestational age, gestational age at delivery                            | AN was associated with higher maternal age, higher maternal education, multiparity, and preterm birth | The rate of AN decreased with maternal overweight and obesity in a dose-response manner | 5              |

AN: anorexia nervosa, BN: bulimia nervosa, HC: healthy control, ED: eating disorder, EDI-2: Eating Disorder Inventory-2, EAT-26: Eating Attitude Test-26, BITE: Bulimic Investigatory Test, FSS: female same-sex twins, FOS: female opposite-sex twins, MOS: male opposite-sex twins, MSS: male same-sex twins, SS: same sex, OS: opposite sex, VPT: very preterm, BMI: body mass index
Culbert et al. 2008, 2013; Raevuori et al. 2008; Baker et al. 2009; Nicholls and Viner 2009; Smith et al. 2010; Wehkalampi et al. 2010; Coombs et al. 2011; Quinton et al. 2011; Vellisca et al. 2013; Micali et al. 2015; Sacks et al. 2016). MMAT scores were lower for cross-sectional and case-control (ranging from 2 to 4 points) than cohort studies (ranging from 3 to 5 points).

Notwithstanding the aforementioned weaknesses, some data gathered by the most robust studies should be acknowledged (see Table 2): in fact, mixed diagnoses of EDs were associated with maternal stress during pregnancy and preterm birth; still, BN was consistently associated with maternal psychosocial stress during pregnancy. Finally, multiple factors, according to the available data, resulted to be related to the onset of AN in the offspring: higher maternal age, preeclampsia and eclampsia, multiparity, hypoxic complications, preaturity or preterm birth (< 32 weeks), and being small for gestational birth size.

Maternal factors

Maternal age (16 studies)

A total of 16 studies assessed maternal age as a possible factor linked to the risk for AN or BN (Cnattingius et al. 1999; Feingold et al. 2002; Lindberg and Hjern 2003; Montgomery et al. 2005; Favaro et al. 2006, 2008; Wehkalampi et al. 2010; Nosarti et al. 2012; Allen et al. 2016).

Table 2 Results summary of the studies included in this review grouped by diagnoses of eating disorders

| Eating disorders (EDs) | Anorexia nervosa (AN) | Bulimia nervosa (BN) |
|------------------------|-----------------------|----------------------|
| Factors supported by more robust evidence | Maternal stress during pregnancy | Higher maternal age | Maternal stress during pregnancy |
| | Preterm birth | Preeclampsia and eclampsia | |
| | | Multiparity | |
| | | Hypoxic complications | |
| | | Prematurity or preterm birth (< 32 weeks) | |
| | | Small for gestational age or lower birth size | |
| Factors supported by less robust evidence | Gestational diabetes | Low maternal weight | Maternal smoke habit |
| | Vit. D deficiency | Viral infection during pregnancy | Vit. D deficiency |
| | | Season of birth (spring) | Dysmaturity signs |
2013; Goodman et al. 2014; Romero-Martínez and Moya-Albiol 2014; Tenconi et al. 2015; St-Hilaire et al. 2015; Lofrano-Prado et al. 2015; Sacks et al. 2016; Razaz and Cnattingius 2018). Five studies out of 16 found that higher maternal age was significantly associated with an increased risk of AN (Cnattingius et al. 1999; Lindberg and Hjern 2003; Goodman et al. 2014; Tenconi et al. 2015; Razaz and Cnattingius 2018). A register study found instead that adolescents with mothers who were young at the time of birth had a lower risk of developing AN compared with the adolescents of mothers who were 25–28 years old at birth (Lindberg and Hjern 2003).

Maternal weight (7 studies)

Out of a total of 7 studies addressing maternal weight (Montgomery et al. 2005; Wehkalampi et al. 2010; Allen et al. 2013; Romero-Martínez and Moya-Albiol 2014; Matinolli et al. 2016; Sacks et al. 2016; Razaz and Cnattingius 2018), the majority did not show significant associations between maternal body mass index (BMI) and the onset of either AN or BN in the offspring. In contrast, a recent study found that the risks of AN in girls born at term decreased with maternal overweight and obesity in a dose-response manner (Razaz and Cnattingius 2018).

Maternal cigarette smoking (2 studies) and alcohol drinking (7 studies)

Only 2 studies investigated maternal alcohol use and both failed to show any associations between maternal alcohol use and EDs onset in the offspring (Feingold et al. 2002; Allen et al. 2013). Seven studies investigated instead the role of maternal cigarette smoke (Montgomery et al. 2005; Nicholls and Viner 2009; Wehkalampi et al. 2010; Allen et al. 2013; Goodman et al. 2014; Matinolli et al. 2016; Razaz and Cnattingius 2018), reporting an association between maternal smoking and BN diagnosis in the offspring, even after controlling for confounding factors such as offspring BMI in adulthood or variation between childhood and adult BMI (Montgomery et al. 2005). In contrast, another study found a strong negative association between mother’s smoking and AN, but this was substantially attenuated upon adjustment for parental education (Goodman et al. 2014). Still, all the other studies did not identify an association (Nicholls and Viner 2009; Wehkalampi et al. 2010; Allen et al. 2013; Matinolli et al. 2016; Razaz and Cnattingius 2018).

Maternal stress during pregnancy (3 studies)

Only 3 studies assessed maternal stress during pregnancy and EDs diagnosis in the offspring and all consistently found a significant association (Taborelli et al. 2013; St-Hilaire et al. 2015; Su et al. 2016). A large population-based cohort study found that girls who were born from mothers who lost a close relative from 1 year before the beginning of pregnancy to the whole duration of pregnancy had an increased risk of suffering from an EDs than healthy controls, with similar results for mixed EDs and BN, but not for AN (Su et al. 2016). Similarly, it has been observed that the daughters of mothers with chronic anxiety during pregnancy had an increased risk of AN (Taborelli et al. 2013). Additionally, maternal stress in the third trimester of pregnancy was associated with elevated scores on a screening tool for EDs (St-Hilaire et al. 2015).

Pregnancy complications

Preeclampsia, eclampsia, pregnancy hypertension (9 studies), and maternal anemia (3 studies)

Nine studies investigated preeclampsia, eclampsia, pregnancy hypertension (Cnattingius et al. 1999; Foley et al. 2001; Feingold et al. 2002; Lindberg and Hjern 2003; Favaro et al. 2006, 2008; Tenconi et al. 2015; Matinolli et al. 2016), and three maternal anemia (Favaro et al. 2006; Nicholls and Viner 2009; Tenconi et al. 2015). Out of 9 studies, only 3 of them found that AN was significantly associated with preeclampsia (Lindberg and Hjern 2003; Favaro et al. 2006; Tenconi et al. 2015), while only in one work (Favaro et al. 2006) an association with maternal anemia during pregnancy was found. Nevertheless, the same group disconfirmed such an association in a more recent study (Tenconi et al. 2015).

Gestational diabetes (7 studies)

Seven studies assessed the role of gestational diabetes (Cnattingius et al. 1999; Favaro et al. 2006, 2008; Nicholls and Viner 2009; Tenconi et al. 2015; Matinolli et al. 2016; Sacks et al. 2016). Out of the available studies, only 2 found an association between gestational diabetes and EDs in the offspring, the first on a sample with mixed ED diagnoses (Sacks et al. 2016) and the latter on a sample with AN (Favaro et al. 2006). However, a more recent study did not confirm this datum (Tenconi et al. 2015).

Maternal viral infection (2 studies)

Two studies analyzed the association between in utero viral infection exposition, in particular rubella (Foley et al. 2001; Favaro et al. 2011) and chickenpox (Favaro et al. 2011). Only one study found that exposure to rubella or chickenpox during the sixth month of pregnancy was associated with an increased risk of developing AN in the offspring (Favaro et al. 2011).
Vitamin D deficiency (1 study)

One study assessed the role of vitamin D deficiency in pregnancy, finding that EDs were predicted by low maternal vitamin D level at 18-week pregnancy, even after controlling for family sociodemographic factors, BMI, and depressive symptoms (Allen et al. 2013). However, BN was the only disorder in which the risk remained significantly increased when EDs were assessed separately.

Effect of sex hormones (12 studies)

Six studies assessed prenatal androgen exposure using the 2D:4D ratio (Klump et al. 2006; Smith et al. 2010; Coombs et al. 2011; Quinton et al. 2011; Romero-Martínez and Moya-Albiol 2014; Culbert et al. 2015). In general, the 4th digit tends to be longer than the 2nd in males, whereas in females the 2nd and 4th digits tend to be of equal length (Berenbaum et al. 2009). Lower 2D:4D ratios (second finger shorter than the fourth finger) point to higher prenatal androgen exposure (Berenbaum et al. 2009). Two studies (Klump et al. 2006; Culbert et al. 2015) found that ED symptomatology [i.e., body dissatisfaction (Klump et al. 2006), weight preoccupation (Klump et al. 2006), compensatory behaviors (Klump et al. 2006), and disordered eating (Culbert et al. 2015)] was associated with lower levels of prenatal testosterone. In males, greater prenatal testosterone exposure is associated with less disordered eating, less drive for leanness, but increased drive for muscularity (Smith et al. 2010). Two other studies found that AN was instead significantly associated with higher testosterone exposure during pregnancy (Quinton et al. 2011; Romero-Martínez and Moya-Albiol 2014), while BN was associated with lower levels of prenatal testosterone (Quinton et al. 2011).

Multiparity (5 studies)

Out of 5 studies assessing the role of multiparity in the risk of EDs onset (Cnattingius et al. 1999; Feingold et al. 2002; Nosarti et al. 2012; Goodman et al. 2014; Razaz and Cnattingius 2018), only 2 of them found an association with later offspring diagnosis of AN (Goodman et al. 2014; Razaz and Cnattingius 2018).

Vaginal bleeding (7 studies)

Despite 7 studies assessing the role of vaginal bleeding during pregnancy, none of them found an association with offspring ED diagnosis/symptomatology (Cnattingius et al. 1999; Shoebridge and Gowers 2000; Foley et al. 2001; Feingold et al. 2002; Favaro et al. 2006, 2008; Tenconi et al. 2015).

Obstetric complications

Three studies found that an increasing number of obstetric complications was correlated to a higher risk of EDs in the offspring, specifically AN (Foley et al. 2001; Favaro et al. 2006) and BN (Foley et al. 2001; Favaro et al. 2006; Lofrano-Prado et al. 2015), and a lower age of onset of the EDs (Favaro et al. 2006). Few obstetric complications have been associated with BN, but only one study found an association between bulimic symptoms in college students and the presence of any obstetric complications at birth (Lofrano-Prado et al. 2015).

Hypoxic complications (7 studies)

Seven studies assessed hypoxic complications, namely, umbilical cord wrapped around the neck, need for O2, and placental infarction (Foley et al. 2001; Feingold et al. 2002; Lindberg and Hjern 2003; Favaro et al. 2006, 2008; Nicholls and Viner 2009; Tenconi et al. 2015). As a result, only 3 studies found an association between hypoxic complications and AN (Lindberg and Hjern 2003; Favaro et al. 2006; Tenconi et al. 2015) or BN (Favaro et al. 2006).

Breech delivery (5 studies)

Out of 5 studies assessing breech delivery (Foley et al. 2001; Feingold et al. 2002; Lindberg and Hjern 2003; Favaro et al. 2006; Tenconi et al. 2015), only one study (Lindberg and Hjern 2003) found an association with later AN diagnosis.

Cephalohematoma (6 studies)

Cephalohematoma was investigated by 6 studies (Cnattingius et al. 1999; Lindberg and Hjern 2003; Favaro et al. 2006, 2008; Goodman et al. 2014; Tenconi et al. 2015) but only 2
of them found an association with AN diagnosis in the offspring (Cnattingius et al. 1999; Lindberg and Hjern 2003).

Premature rupture of the membranes (8 studies)

Eight studies assessed premature rupture of the membranes (Cnattingius et al. 1999; Foley et al. 2001; Feingold et al. 2002; Lindberg and Hjern 2003; Favaro et al. 2006, 2008; Goodman et al. 2014; Tenconi et al. 2015). Only one study found an association with AN (Lindberg and Hjern 2003) but was then disconfirmed by a later larger work (Goodman et al. 2014).

Cesarean section delivery (8 studies) and forceps or vacuum delivery (8 studies)

Eight studies investigated the role of cesarean section delivery (Cnattingius et al. 1999; Shoebridge and Gowers 2000; Foley et al. 2001; Feingold et al. 2002; Goodman et al. 2014; Micali et al. 2015; Lofrano-Prado et al. 2015; Razaz and Cnattingius 2018) and forceps or vacuum delivery (Cnattingius et al. 1999; Shoebridge and Gowers 2000; Foley et al. 2001; Favaro et al. 2006, 2008; Goodman et al. 2014; Tenconi et al. 2015; Razaz and Cnattingius 2018). Only one study found a weak evidence of an association with AN diagnosis (Goodman et al. 2014) while another (Micali et al. 2015) found that cesarean section was instead associated with compensatory behaviors.

Neonatal factors

Gestational age, birth size, and size for gestational age (18 studies)

Eighteen studies assessed prematurity in terms of gestational age, birth size, or birth size for gestational age (Cnattingius et al. 1999; Shoebridge and Gowers 2000; Foley et al. 2001; Feingold et al. 2002; Lindberg and Hjern 2003; Favaro et al. 2006, 2008; Nicholls and Viner 2009; Wehkalampi et al. 2010; Nosarti et al. 2012; Allen et al. 2013; Goodman et al. 2014; Micali et al. 2015; Tenconi et al. 2015; St-Hilaire et al. 2015; Matinolli et al. 2016; Sacks et al. 2016; Razaz and Cnattingius 2018).

Gestational age (< 32 weeks) has been consistently found to be associated with EDs, even adjusting for confounders (Nosarti et al. 2012; Foley et al. 2001; Goodman et al. 2014; Cnattingius et al. 1999; Lindberg and Hjern 2003; Razaz and Cnattingius 2018). With more detail, low gestational age was associated with later AN diagnosis with an odds ratio ranging from 1.9 (95% CI 1.2–3.3; Lindberg and Hjern 2003) to 3.2 (95% CI 1.6–6.2; Cnattingius et al. 1999).

Three studies found that AN diagnosis was associated with being small for gestational age at birth (Cnattingius et al. 1999) or having a Ponderal Index < 25 (Favaro et al. 2006; Tenconi et al. 2015).

Concerning BN, two studies found that BN was associated with low birth weight for gestational age (Favaro et al. 2006; Tenconi et al. 2015); however, another study reported the opposite result (Goodman et al. 2014).

Two studies on unaffected individuals found that adolescents and young adults born preterm showed lower scores on eating psychopathology (i.e., drive for thinness, body dissatisfaction, and bulimia) than those who were not born preterm (Wehkalampi et al. 2010) even after controlling for confounders (Matinolli et al. 2016).

Apgar score or dysmaturity signs (7 studies)

Seven studies assessed Apgar score (Cnattingius et al. 1999; Shoebridge and Gowers 2000; Lindberg and Hjern 2003; Nosarti et al. 2012) or dysmaturity signs (Favaro et al. 2006, 2008; Tenconi et al. 2015) such as hypotonia, hyporeactivity, hypothermia, tremors, and feeding problems at birth. One study found that AN diagnosis was associated with low Apgar score at birth (Lindberg and Hjern 2003); in keeping with these findings, the other two studies found that hyporeactivity was a significant independent predictor of the development of AN even after adjusting for confounders (Favaro et al. 2006; Tenconi et al. 2015). Additionally, the same studies found neonatal hyporeactivity and early eating difficulties as adjusted risk factors for BN (Favaro et al. 2006; Tenconi et al. 2015). In another study, the presence of signs of neonatal dysmaturity influenced the development of high harm avoidance, a risk factor of EDs (Favaro et al. 2008).

Season of birth (5 studies)

Five studies assessed season or month of birth and the association with EDs providing consistent support to an association of AN with being born in spring (Morgan and Lacey 2000; Favaro et al. 2011; Vellisca et al. 2013; Winje et al. 2013; Allen et al. 2013).

Discussion

The aim of this systematic review was to highlight the association between prenatal and perinatal factors and the subsequent development of EDs, investigating the hypothesis that these factors could impair neurodevelopment, similarly to the model proposed for schizophrenia (Geddes and Lawrie 1995; Verdoux et al. 1997; Geddes et al. 1999; Cannon et al. 2000; Clarke et al. 2011; Rapoport et al. 2012). When analyzing the main findings of this review, it should be also borne in mind that it has been brought to the surface the inconsistency of the available body of evidence on this topic and the lack of a
robust framework able to explain the possible relationships between prenatal and perinatal factors and clinical variables in EDs. Nevertheless, some relevant main findings emerged as well: first, maternal stress during pregnancy and preterm birth emerged as the most supported factors impacting on a diagnosis of EDs in the offspring. Similarly, maternal stress during pregnancy was robustly associated with the onset of BN. Finally, the association between prenatal and perinatal factors and AN resulted to be complex and many-sided: in fact, higher maternal age, preeclampsia and eclampsia, multiparity, hypoxic complications, and prematurity or preterm birth (<32 weeks) or being small for gestational age or with a low birth size were the most sound factors in the association with the onset of AN in the offspring.

That said, in keeping with the second aim of this work, future lines of research can be outlined in order to bridge these gaps in the literature: for example, no studies investigated binge eating disorder (BED) and further works may want to tackle the aforementioned methodological weaknesses (i.e., composite variable for prenatal and perinatal risk factors; lack of a shared definition of such risk factors) with ad hoc study designs. As a first step, as suggested by literature (Krug et al. 2013), all factors should be divided into four groups: pregnancy complications, labor and delivery complications, and fetal distress signs/neonatal complications. As a second step, other candidates proposed by this review could be added as well, including: mothers’ characteristics (i.e., age, smoke habit, weight, stressful events during pregnancy) and timing (seasonality). Doing so, it will be finally possible to conduct a meta-analysis of these data yielding quantitative results as well.

With more detail, the main findings for each category (i.e., maternal factors, pregnancy complications, obstetric complications, and neonatal factors) are described below.

Maternal factors

With respect to maternal factors, maternal age has been deepened by a number of robust studies finding that a higher maternal age was significantly associated with an increased risk of AN in the offspring (Cnattingius et al. 1999; Feingold et al. 2002; Favaro et al. 2011; Goodman et al. 2014; Razaz and Cnattingius 2018). However, Lofrano-Prado and collaborators (Lofrano-Prado et al. 2015) found a contrasting result, which could be due both to a greater risk of pregnancy complications in older mothers or to later socio-educational and environmental factors linked to older mothers, so future studies are needed to clarify these matters.

Recent evidence indicates that another maternal factor, namely, obesity and metabolic diseases, may have a long-term impact on psychiatric conditions of the offspring, such as attention deficit hyperactive disorder, autism, and schizophrenia (Rivera et al. 2015). In contrast, in the field of EDs, only one study found that the risk of AN in girls born at term decreased with maternal overweight and obesity in a dose-response manner (Razaz and Cnattingius 2018). However, such a finding was not confirmed in the sibling control analyses, so other genetic or familial environmental factors may be involved as well.

Out of seven studies available, maternal smoke was reported as significant only by two studies with contrasting results (Montgomery et al. 2005; Goodman et al. 2014). In fact, on one hand, a positive association with BN, even after controlling for confounding factors such as offspring BMI in adulthood or variation between childhood and adult BMI, was reported (Montgomery et al. 2005), but on the other hand, a negative association between mother’s smoking and AN was shown, even if attenuated upon adjustment for parental education (Goodman et al. 2014). Although a larger number of studies (i.e., 7) investigated the association between maternal alcohol use and EDs onset in the offspring, no significant data emerged; since this datum is not in line with other fields of psychiatry (Pagnin et al. 2019), future studies are needed to clarify this issue.

Interestingly, psychosocial stress was found to be strongly associated with both EDs and BN in the offspring, on the basis of all the available studies (i.e., 3) that supported this datum with robust evidence (Taborelli et al. 2013; St-Hilaire et al. 2015; Su et al. 2016). Also, maternal stress has been reported to entail a greater risk of impulsive and compensatory behaviors, as hypothesized also in other fields of psychiatry (Abbott et al. 2018). This line of research needs a deeper investigation because it could underlie an important link between endocrine functions, genetics, and temperamental traits. Stress activates the HPA axis, increasing the release of glucocorticoids (St-Hilaire et al. 2015) that could cross the maternal placenta and impact on the development of metabolism, fetal growth, and immune functions during pregnancy. Glucocorticoids could also affect the development of the fetal HPA axis, causing an alteration in stress-response mechanisms, emotional dysregulation, and increased risk for anxiety disorders and EDs in childhood and adulthood (Meyer and Hamel 2014; St-Hilaire et al. 2015).

Pregnancy complications

Concerning pregnancy complications, gestational diabetes could alter fetal neurodevelopment during critical periods.
exposing the fetus to elevated glucose levels (Georgieff 2006). Out of seven studies, two found an association between gestational diabetes and EDs in the offspring (Favaro et al. 2006; Sacks et al. 2016), although subsequent work partially disconfirmed such findings (Tenconi et al. 2015). Similarly, more research is needed also on the role of viral infection during pregnancy: only the study by Favaro et al. (2011) reported that exposure to rubella or chickenpox during the sixth month was associated with an increased risk of developing AN in the offspring.

Animal studies suggested that transient prenatal vitamin D deficiency is associated with altered brain development (Ali et al. 2018) and low maternal vitamin D during pregnancy was identified as a significant predictor of later schizophrenia (McGrath et al. 2010, 2011) and autism (Ali et al. 2018) in the offspring. Surprisingly, only one study was conducted in the ED field, reporting data in line with those of general psychiatry: in fact, EDs were predicted by low maternal vitamin D level at 18-week pregnancy, even after controlling for family sociodemographic factors, BMI, and depressive symptoms (Allen et al. 2013). However, BN was the only disorder in which the risk remained significantly increased when EDs were assessed separately, so no definitive conclusions can be drawn.

Exposure to testosterone during pregnancy has been associated with organizational permanent effects of eating behavior: in animal models, prenatal testosterone exposure increased food intake in male mammals, while in females low levels of testosterone were associated with later restrictive eating behavior (Donohoe and Stevens 1983; Madrid et al. 1993). Lower levels of prenatal testosterone exposure have been associated with body dissatisfaction, weight preoccupation, binge eating, and compensatory behaviors in females (Klump et al. 2006). It has been also suggested that the relatively low level of testosterone before birth in females permits their brains to respond to estrogens at puberty when the hormones activate the genes contributing to disordered eating in vulnerable girls (Klump et al. 2006). Prenatal exposure to male hormones could be indirectly investigated in adult females using the finger-length ratios (2D:4D), a sexually dimorphic trait that correlates with prenatal androgen exposure (Klump et al. 2006) or in opposite-sex twins cohorts, where the female fetus is exposed to higher levels of testosterone by sharing the womb with the male fetus (Resnick et al. 1993; Cohen-Bendahan et al. 2004, 2005). The hypothesis that higher prenatal testosterone exposure could increase food intake and protect against the development of disordered eating symptoms is intriguing although debated. Studies of 2D:4D ratios have yielded more positive evidence than those examining females from opposite-sex twin pairs, but the overall results are very controversial. Future studies should focus on direct assessment of sex hormone levels during pregnancy and perform a better confounder analysis on the other risk factors.

**Obstetric complications**

Obstetric complications that seem to have more robust evidence of association with later AN onset are hypoxic complications (Lindberg and Hjern 2003; Favaro et al. 2006; Tenconi et al. 2015), breech delivery (Lindberg and Hjern 2003), and cephalohematoma (Cnattingius et al. 1999; Lindberg and Hjern 2003). Preeclampsia and eclampsia, severe hypoxic and fetal hypoperfusion pregnancy complications, were associated with a later diagnosis of AN in three more robust-evidence cohort studies, two from the same expanded pool of patients (Lindberg and Hjern 2003; Favaro et al. 2006; Tenconi et al. 2015). Observation of neuropsychological deficits (Galdersisi et al. 2003), subtle neurological abnormalities (Gillberg et al. 1994), and nonreversible morphological brain changes (Katzman et al. 1997; Chowdhury et al. 2003) might suggest that impairment in neurodevelopment could be one of the possible pathways for the development of an ED (Connan et al. 2003). Obstetric complications might have more than one role in their etiopathogenesis; they could cause hypoxia-induced damage to the brain thus impairing the neurodevelopment of the fetus (Cannon et al. 2000). Also, the adequacy of nutrition during pregnancy and the postnatal period could influence the adults’ nutritional status and their appetite programming throughout life (Jones et al. 2017). Perinatal hypoxia/ischemia could cause disturbances of the dopaminergic system that persists in adulthood and impairs the neurotrophic signaling critical for pre- and postnatal brain development (Giannopoulou et al. 2018).

Despite eight studies investigating the role of cesarean section delivery, only weak evidence is available on the association with AN diagnosis (Goodman et al. 2014) or ED symptomatology (Micali et al. 2015). Similarly, also data on the premature rupture of the membranes garnered an overall weak association with EDs.

**Neonatal factors**

Prematurity, in particular very preterm birth (≤32 weeks), or being small for gestational age and birth size have been associated with several cohort studies to AN (Cnattingius et al. 1999; Foley et al. 2001; Lindberg and Hjern 2003; Favaro et al. 2006; Goodman et al. 2014; Razaz and Cnattingius 2018) and EDs (Nosarti et al. 2012). Interestingly, a clear dose-response pattern has been found between lower gestational age and higher risk for AN, with a gradient observed even within the term (births at 37 weeks of gestation) (Goodman et al. 2014). Cnattingius et al. (1999) showed that among girls born very preterm, the risk of subsequent development of AN was higher among girls who were small for
gestational age (OR 5.7, 95% CI 1.1–28.7) than among girls with higher birth weight for gestational age (OR 2.7, 95% CI 1.2–5.8). Less clear is the association between BN and birth weight: two cohort studies (Favaro et al. 2006; Tenconi et al. 2015) from the same population found that BN was associated with being born small for gestational age; however, another study found the opposite result (Goodman et al. 2014). Micali et al. (2015) despite not having found any associations between perinatal predictors and ED psychopathology, observed that those very preterm adults that at the age of 21 years presented with ED symptoms had a smaller gray matter volume in the posterior cerebellum and a smaller white matter volume in the fusiform gyrus bilaterally at the age of 14–15 years. Early alteration of the cerebellum sub-networks linked with somatosensory, interoceptive, and emotional processings was recently found (Gaudio et al. 2018). These findings, if confirmed by further studies, could help to explain the abnormal integration of somatosensory and homeostatic signals, which may lead to body image disturbances in AN.

Broadly speaking, premature newborns need to face a difficult environment that increases stress potentially influencing the psychic and brain development in turn generating epigenetic changes. Follow-up studies often report neurocognitive inabilities with multi-level minimal impairments (Fumagalli et al. 2018; Nist et al. 2019). In the same vein, predisposing factors for EDs could impact not only directly on the relationship with food but also indirectly (i.e., socio-emotional and/or neuropsychological difficulties) increasing individuals’ vulnerability to factors occurring later in the lifespan, for example, cultural factors and hormonal changes in adolescence.

The findings that healthy individuals born preterm scored lower on eating psychopathology than HC’s born at term (Wekkalampi et al. 2010; Matinolli et al. 2016) are apparently in contrast with some aforementioned studies. Notwithstanding, these inconsistencies could be due to a sub-optimal diagnostic assessment (EDI-2 and not the ICD/DSM gold standard) or to a sample selection bias or to the lack of adjustment for confounders. Future studies on healthy individuals born preterm are warranted to better understand their eating style.

Interestingly, dysmaturity signs (e.g., hyporeactivity) have been linked to personality alterations in AN (Favaro et al. 2006; Tenconi et al. 2015) and BN (Favaro et al. 2006; Tenconi et al. 2015). As shown by Favaro et al. (2008), neonatal dysmaturity influences the development of particular temperamental dimensions (Cloninger et al. 1993), such as harm avoidance (HA), a temperament dimension that has been associated with a higher risk of AN onset (Atiyeh et al. 2015). HA reflects the tendency to respond intensely to aversive stimuli and involves anticipatory worry about possible problems (Favaro et al. 2008). It is considered a marker of emotional vulnerability to depression (Kampman and Poutanen 2011) not only because individuals with high HA are more anxiety-prone but also because of their limited ability to recover from depression.

Seasonality is still a controversial topic that could underlie many other factors, such as gestational vitamin D, exposure to infectious agents, temperature and weather, and/or pregnancy and birth complications, all of which have the potential to influence fetal or infant neurodevelopment. Results are mixed, with studies supporting the season of birth hypothesis for generic EDs (Eagles et al. 2001; Watkins et al. 2002; Waller et al. 2002), although this association is lost when both AN (Button and Aldridge 2007; Vellisca et al. 2013; Winje et al. 2013) and BN (Morgan and Lacey 2000; Button and Aldridge 2007) are studied separately. Given the several methodological issues affecting these data (e.g., different latitudes, the accuracy of diagnosis, sample sizes, and comparison groups), caution is required when reading these findings.

Conclusions

The investigation of the association of prenatal and perinatal factors with later onset of psychiatric conditions is particularly hard to perform from a methodological standpoint. This is even more true when such an evaluation is applied to quite rare conditions like AN (Hoek 2006). Notwithstanding, the available body of evidence supports maternal stress during pregnancy and preterm birth as associated with the development of mixed diagnoses of EDs; still, BN was consistently associated with maternal psychosocial stress during pregnancy. Finally, multiple factors were reported to have an impact on the onset of AN in the offspring, namely, higher maternal age, preeclampsia and eclampsia, multiparity, hypoxic complications, prematurity, or preterm birth (<32 weeks), and being small for gestational birth size. However, our review contributed also to shed light on the plethora of methodological inconsistencies and flaws that characterize these lines of research (e.g., inclusion biases, self-report assessments of the EDs, lack of a shared definition of prenatal/perinatal factors, to name a few), potentially generating new ideas on how to tackle these weaknesses and finally provide a meta-analysis on this clinically relevant topic. More studies on larger samples evaluating multiple factors with a longitudinal design—the most fruitful research methodology given the research questions—are necessary to draw more solid conclusions on this topic. Also, as reported earlier (Krug et al., 2013), a shared definition of risk factors is required; in this light, the classification system adopted in this review (i.e., maternal factors, pregnancy complications, obstetric complications, and neonatal factors) could be a starting point to promote the additional debate on prenatal and perinatal factors in the field of EDs in turn encouraging well-designed studies.
References

Abbott PW, Gumusoglu SB, Bitte J, Beverdorp DQ, Stevens HE (2018) Prenatal stress and genetic risk: how prenatal stress interacts with genetics to alter risk for psychiatric illness. Psychoneuroendocrinology 90:9–21. https://doi.org/10.1016/j.psyneuen.2018.01.019

Ali A, Cui X, Eyles D (2018) Developmental vitamin D deficiency and autism: putative pathogenic mechanisms. J Steroid Biochem Mol Biol 175:108–118. https://doi.org/10.1016/j.jsbmb.2016.12.018

Allen KL, Byrne SM, Kusel MMH, Hart PH, Whitehouse AJO (2013) Maternal vitamin D levels during pregnancy and offspring eating disorder risk in adolescence. Int J Eat Disord 46:669–676. https://doi.org/10.1002/eat.22147

Atiyeh M, Miettunen J, Raevuori-Helkamaa A (2015) A meta-analysis of temperament in eating disorders. Eur Eat Disord Rev 23:89–99. https://doi.org/10.1002/eat.22342

Baker JH, Lichtenstein P, Kendler KS (2009) Intrauterine testosterone exposure and risk for disordered eating. Br J Psychiatry J Ment Sci 194:375–376. https://doi.org/10.1192/bjp.bp.108.054692

Baker JH, Schaumberg K, Monn-Chernoff MA (2017) Genetics of anorexia nervosa. Curr Psychiatry Rep 19:84. https://doi.org/10.1007/s11920-017-0842-2

Berenbaum SA, Bryk KK, Nowak N, Quigley CA, Moffat S (2009) Fingers as a marker of prenatal androgen exposure. Endocrinology 150:5119–5124. https://doi.org/10.1210/en.2009-0774

Button E, Aldridge S (2007) Season of birth and eating disorders: patterns across diagnoses in a specialized eating disorders service. Int J Eat Disord 40:468–471

Cannon TD, Rosso IM, Hollister JM, Beanrden CE, Sanchez LE, Hadley T (2000) A prospective cohort study of genetic and perinatal influences in the etiology of schizophrenia. Schizophr Bull 26:351–366

Chowdhury U, Gordon I, Lask B, Watkins B, Watth H, Christie D (2003) Early-onset anorexia nervosa: is there evidence of limbic system imbalance? Int J Eat Disord 33:388–396. https://doi.org/10.1002/eat.10155

Clarke MC, Tanskanen A, Huttunen M, Leon DA, Murray RM, Jones PB, Cannon M (2011) Increased risk of schizophrenia from additive interaction between infant motor developmental delay and obstetric complications: evidence from a population-based longitudinal study. Am J Psychiatry 168:1295–1302. https://doi.org/10.1176/appi.ajp.2011.11010011

Cloninger CR, Svrakic DM, Przybeck TR (1993) A psychobiological model of temperament and character. Arch Gen Psychiatry 50:975–990

Chattingius S, Hultman CM, Dahl M, Sparén P (1999) Very preterm birth, birth trauma, and the risk of anorexia nervosa among girls. Arch Gen Psychiatry 56:634–638

Cohen-Bendahan CCC, Buitelaar JK, van Gooszen SHM, Cohen-Kettenis PT (2004) Prenatal exposure to testosterone and functional cerebellar lateralization: a study in same-sex and opposite-sex twin girls. Psychoneuroendocrinology 29:911–916. https://doi.org/10.1016/j.psyneuen.2003.07.001

Cohen-Bendahan CCC, Buitelaar JK, van Gooszen SHM, Orlebeke JF, Cohen-Kettenis PT (2005) Is there an effect of prenatal testosterone on aggression and other behavioral traits? A study comparing same-sex and opposite-sex twin girls. Horm Behav 47:230–237. https://doi.org/10.1016/j.yhbeh.2004.10.006

Conna F, Campbell IC, Katzman M, Lightman SL, Treasure J (2003) A neurodevelopmental model for anorexia nervosa. Psychol Behav 79:13–24

Coombs E, Brosnan M, Bryant-Waugh R, Skevington SM (2011) An investigation into the relationship between eating disorder psychopathology and autistic symptomatology in a non-clinical sample. Br J Clin Psychol 50:326–338. https://doi.org/10.1348/014466510X524408

Culbert KM, Breedlove SM, Burt SA, Klump KL (2008) Prenatal hormone exposure and risk for eating disorders: a comparison of opposite-sex and same-sex twins. Arch Gen Psychiatry 65:329–336. https://doi.org/10.1001/archgenpsychiatry.2007.47

Culbert KM, Breedlove SM, Siick CL, Burt SA, Klump KL (2013) The emergence of sex differences in risk for disordered eating attitudes during puberty: a role for prenatal testosterone exposure. J Abnorm Psychol 122:420–432. https://doi.org/10.1037/a0031791

Culbert KM, Breedlove SM, Siick CL, Keel PK, Neale MC, Boker SM, Burt SA, Klump KL (2015) Age differences in prenatal testosterone’s protective effects on disordered eating symptoms: developmental windows of expression? Behav Neurosci 129:18–36. https://doi.org/10.1037/bne0000034

Dalle Grave R (2011) Eating disorders: progress and challenges. Eur J Intern Med 22:153–160. https://doi.org/10.1016/j.ejim.2010.12.010

Eagles JM, Andrew JE, Johnston MI, Easton EA, Millar HR (2001) Season of birth in females with anorexia nervosa in Northeast Scotland. Int J Eat Disord 30:167–175

Fairburn CG, Harrison PJ (2003) Eating disorders. Lancet Lond Engl 361:407–416. https://doi.org/10.1016/S0140-6736(03)12378-1

Favaro A, Tencini E, Santonastaso P (2006) Perinatal factors and the risk of developing anorexia nervosa and bulimia nervosa. Arch Gen Psychiatry 63:82–88. https://doi.org/10.1001/archpsyc.63.1.82

Favaro A, Tencini E, Santonastaso P (2008) The relationship between obstetric complications and temperament in eating disorders: a mediation hypothesis. Psychosom Med 70:372–377. https://doi.org/10.1097/PSY.0b013e318164604e

Favaro A, Tencini E, Ceschin L, Zanetti T, Bosello R, Santonastaso P (2011) In utero exposure to virus infections and the risk of developing anorexia nervosa. Psychol Med 41:2193–2199. https://doi.org/10.1017/S0033291710002655

Feingold E, Sheir-Neiss G, Melnychuk J, Bachrach S, Paul D (2002) Eating disorder symptomatology is not associated with pregnancy...
and perinatal complications in a cohort of adolescents who were born preterm. Int J Eat Disord 31:202–209
Foley DL, Thacker LR, Aggen SH, Neale MC, Kendler KS (2001) Pregnancy and perinatal complications associated with risks for common psychiatric disorders in a population-based sample of female twins. Am J Med Genet 105:426–431
Fumagalli M, Provenzi L, De Carli P et al (2018) From early stress to 12-month development in very preterm infants: preliminary findings on epigenetic mechanisms and brain growth. PLoS One 13:e0196002. https://doi.org/10.1371/journal.pone.0196002
Galeris S, Mucci A, Montelone P, Sorrentino D, Piegaro G, Maj M (2003) Neurocognitive functioning in subjects with eating disorders: the influence of neuroactive steroids. Biol Psychiatry 53:921–927
Gardener H, Spiegelman D, Buka SL (2009) Prenatal risk factors for autism: comprehensive meta-analysis. Br J Psychiatry J Ment Sci 195:7–14. https://doi.org/10.1192/bjp.bp.108.051672
Gaudio S, Olivo G, Zobel BB, Schiöth HB (2018) Altered cerebellar–insular–parietal–circular subnetwork in adolescents in the earliest stages of anorexia nervosa: a network–based statistic analysis. Transl Psychiatry 8:1–10. https://doi.org/10.1038/s41398-018-0173-z
Geddes JR, Lawrie SM (1995) Obstetric complications and schizophrenia: a meta-analysis. Br J Psychiatry J Ment Sci 167:786–793
Geddes JR, Verduux H, Takei N, Lawrie SM, Bovet P, Eagles JM, Heun R, McCreadie RG, McNeil TF, O'Callaghan E, Stober G, Willinger U, Murray RM (1999) Schizophrenia and complications of pregnancy and labor: an individual patient data meta-analysis. Schizophr Bull 25:413–423
Georgieff MK (2006) The effect of maternal diabetes during pregnancy on the neurodevelopment of offspring. Minn Med 89:44–47
Giannopoulou I, Pagida MA, Briana DD, Panayotacopolou MT (2018) Perinatal hypoxia as a risk factor for psychopathology later in life: the role of dopamine and neurotrophins. Horm Athens Greece 17:25–32. https://doi.org/10.1007/s42000-018-0007-7
Gillberg C, Råstam M, Gillberg IC (1994) Anorexia nervosa: physical health and neurodevelopment at 16 and 21 years. Dev Med Child Neurol 36:567–575
Goodman A, Heshmati A, Malik N, Koupil I (2014) Associations between birth characteristics and eating disorders across the life course: findings from 2 million males and females born in Sweden, 1975–1998. Am J Epidemiol 179:852–863. https://doi.org/10.1093/aje/kvt445
Hoek HW (2006) Incidence, prevalence and mortality of anorexia nervosa and other eating disorders. Curr Opin Psychiatry 19:389–394. https://doi.org/10.1097/01.yco.0000228759.55277.88
Hong Q, Pluye P, Fàbregues S (2018) Mixed methods appraisal tool (MMAT)
Jacobi C, Fittig E, Bryson SW, Wilfley D, Kraemer HC, Taylor CB (2011) Who is really at risk? Identifying risk factors for subthreshold and full syndrome eating disorders in a high-risk sample. Psychol Med 41:1939–1949. https://doi.org/10.1017/S0033291710002631
Jones C, Pearce B, Barrera I, Mummert A (2017) Fetal programming and eating disorder risk. J Theor Biol 428:26–33. https://doi.org/10.1016/j.jtbi.2017.05.028
Kampman O, Poutanen O (2011) Can onset and recovery in depression be predicted by temperament? A systematic review and meta-analysis. J Affect Disord 135:20–27. https://doi.org/10.1016/j.jad.2010.12.021
Katzman DK, Zipursky RB, Lambe EK, Mikulis DJ (1997) A longitudinal magnetic resonance imaging study of brain changes in adolescents with anorexia nervosa. Arch Pediatr Adolesc Med 151:793–797
King JA, Frank GKW, Thompson PM, Ehrlich S (2018) Structural neuroimaging of anorexia nervosa: future directions in the quest for mechanisms underlying dynamic alterations. Biol Psychiatry 83:224–234. https://doi.org/10.1016/j.biopsych.2017.08.011
Clump KL, Gobrogge KL, Perkins PS et al (2006) Preliminary evidence that gonadal hormones organize and activate disordered eating. Psychol Med 36:539–546. https://doi.org/10.1017/S0033291705006653
Krug I, Taborelli E, Sallis H, Treasure J, Micali N (2013) A systematic review of obstetric complications as risk factors for eating disorder and a meta-analysis of delivery method and prematurity. Physiol Behav 109:51–62. https://doi.org/10.1016/j.physbeh.2012.11.003
Lindberg L, Hjern A (2003) Risk factors for anorexia nervosa: a national cohort study. Int J Eat Disord 34:397–408. https://doi.org/10.1002/eat.10221
Lindström K, Lagerroos P, Gillberg C, Fernell E (2006) Teenage outcome after being born at term with moderate neonatal encephalopathy. Pediatri Neurol 35:268–274. https://doi.org/10.1016/j.pediatrneurol.2006.05.003
Loftsø-Prado MC, do Prado WL, de Barros MVG et al (2015) Obstetric complications and mother’s age at delivery are predictors of eating disorder symptoms among health science college students. Einstein Sao Paulo Braz 13:525–529. https://doi.org/10.1590/S1679-45082015A0336
Lydecker JA, Pitsesky EM, Mitchell KS, Thornton LM, Kendler KS, Reichborn-Kjennerud T, Lichtenstein P, Bulik CM, Mazzeo SE (2013) Association between co-twin sex and eating disorders in opposite sex twin pairs: evaluations in north American, Norwegian, and Swedish samples. J Psychosom Res 72:73–77. https://doi.org/10.1016/j.jpsychores.2011.05.014
Matinolli H-M, Männistö S, Sipola-Leppänen M, Tikkanmäki M, Heinonen K, Lahti J, Lahti M, Wehkalampi J, Järvelin MR, Andersson S, Lano A, Vartia T, Wolke D, Eriksson JG, Vääräsmäki M, Räikönen K, Kajantie E (2016) Body image and eating behavior in young adults born preterm. Int J Eat Disord 49:572–580. https://doi.org/10.1002/eat.22553
McGrath JJ, Eyles DW, Pedersen CB et al (2010) Neonatal vitamin D status and risk of schizophrenia: a population-based case-control study. Arch Gen Psychiatry 67:889–894. https://doi.org/10.1001/archgenpsychiatry.2010.110
McGrath JJ, Hannan AJ, Gibson G (2011) Decanalization, brain development and risk of schizophrenia. Transl Psychiatry 1:e14. https://doi.org/10.1038/tp.2011.16
Meyer JS, Hamel AF (2014) Models of stress in nonhuman primates and their relevance for human psychopathology and endocrine dysfunction. ILAR J 55:347–360. https://doi.org/10.1093/ilarilu/023
Micali N, Kothari R, Nam KW, Gioroukou E, Walshe M, Allin M, Rifkin L, Murray RM, Nosarti C (2015) Eating disorder psychopathology, brain structure, neuropsychological correlates and risk mechanisms in very preterm young adults. Eur Eat Disord Rev J Eat Disord Assoc 23:147–155. https://doi.org/10.1002/erv.2346
Moher D, Liberati A, Tetzlaff J, Altman DG, The PRISMA Group (2009) Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med 6:e1000097. https://doi.org/10.1371/journal.pmed.1000097
Montgomery SM, Ehin A, Ekborn A (2005) Smoking during pregnancy and bulimia nervosa in offspring. J Perinat Med 33:206–211. https://doi.org/10.1515/JPM.2005.038
Morgan JF, Lacey JH (2000) Season of birth and bulimia nervosa. Int J Eat Disord 27:452–458
Nicollas DE, Viner RM (2009) Childhood risk factors for lifetime anorexia nervosa by age 30 years in a national birth cohort. J Am Acad Child Adolesc Psychiatry 48:791–799. https://doi.org/10.1097/CHI.0b013e3181ab8b75
Nist MD, Harrison TM, Steward DK (2019) The biological embedding of neonatal stress exposure: a conceptual model describing the mechanisms of stress-induced neurodevelopmental impairment in preterm infants. Res Nurs Health 42:61–71. https://doi.org/10.1002/nur.21923
