ABSTRACT

**Background:** Hyperemesis gravidarum tends to rapidly progress into Wernicke encephalopathy and Korsakoff syndrome and, therefore, needs to be recognized early and managed promptly with targeted multimodal therapies.

**Objective:** This study critically reviewed the relevant literature on clinical perspectives of hyperemesis gravidarum, Wernicke encephalopathy and Korsakoff syndrome. The secondary objective of this study was to improve the awareness, emphasis on early diagnosis and immediate intervention concerning these sequential syndromes in pregnant women across the board.

**Methods:** Electronic searches (since inception-2019) of PubMed/MEDLINE, Google Scholar, OvidSP, Dove Medical Press, ScienceDomain International (SDI) and Hindawi.com were...
conducted using keywords and Boolean Operators. Hundreds of thousands articles were retrieved which were reviewed independently by two authors and finally 144 articles retained that addressed clinical components of these sequential syndromes along with relevance of thiamine deficiency. **Results:** Evidently, vulnerable women in early stage of gestation tend to develop hyperemesis gravidarum characterized by persistent severe pernicious nausea and vomiting that causes Wernicke encephalopathy defined by variable oculomotor disturbances, ataxia, confusion, metabolic disturbances and Korsakoff syndrome linked with gross memory impairment, confabulation and constipation. The women with these conditions need diagnostic evaluation by means of clinical history, relevant laboratory tests, abdominal ultrasound and brain computerized tomography and magnetic resonance imaging. Most patients need emergency admission, prompt treatment with optimal doses of antiemetics, vitamin B1, and followed by fluid replenishment and electrolyte balance with follow up till the end of pregnancy. Successful maternal and fetal outcome of pregnancy depends on multiple determinants including associated systemic diseases. **Conclusion:** Evidently, a variety of etiological and risk factors in pregnant women determine the initiation of hyperemesis gravidarum that subsequently causes Wernicke encephalopathy and Korsakoff syndrome, and each of which needs prompt multimodal treatment in order to reduce maternal morbidity and increase successful outcome of pregnancy. Although clinical literature concerning these sequential syndromes is huge, further studies are needed to understand their underlying pathophysiological pathways across the world.

**Keywords:** Hyperemesis gravidarum; wernicke encephalopathy and Korsakoff syndrome; thiamine deficiency.

1. **INTRODUCTION**

Pregnancy is a normal physiological process and ends by ninth month with a healthy baby. However, pregnancy may associate with morning sickness, nausea and vomiting of pregnancy (NVP), hyperemesis gravidarum (HG), Wernicke encephalopathy (WE) and Wernicke-Korsakoff syndrome (WKS) [1]. HG is an emergency condition characterized by severe, intractable nausea and vomiting affecting 0.3-3% of all pregnancies [1-3]. NVP is a common condition presenting in 50% to 90% of all gravidas [4,5]. Both recurrent conditions lie on the opposite poles of a spectrum but need suitable interventions. The onset of NVP starts at 4-8 weeks and subsides in 90% cases by 16-20 weeks; however, NVP may persist beyond 20 weeks in 13% of cases [2-7]. Evidently, HG more severe than NVP is typically characterized by pernicious nausea and vomiting (3 times/day) that results in dehydration, electrolyte and fluid imbalance, weight loss (≥5%), ketonuria, physical and psychological debilitation and very often necessitates hospital admission [2,6]. In the past, hundreds of case reports, many studies and reviews mainly from Western world had described HG co-occurring with and complicated by diverse systemic diseases, infections and medications [8-39] (Table 1). A recent systematic review of HG revealed 146 case studies that reported on 177 cases of HG and WE, linked with worsening fetal and maternal outcome [31,40-43]. In specific terms, HG primarily induced by thiamine deficiency is associated with fetal low birth weight, intrauterine growth restriction, preterm delivery, neurodevelopmental anomalies and poor maternal health which adversely impact their future life trajectories [44-48]. In addition, thiamine deficiency in infant-mother domain is a neglected problem worldwide with possible linkage to sudden infant death syndrome [49]. Interestingly, thiamine deficiency may also cause aforesaid fetal problems in pregnant women even without HG [50]. Some reports suggested that the perinatal outcome of pregnant women who suffered from HG remain unaffected [51]. HG often results in an acute thiamine deficiency which is characterized by loss of appetite, nausea and vomiting, fatigue, weakness, anxiety, difficulty in concentration, confusion, disorientation, and memory loss and, if not treated, progresses to or causes WE, KS and Wernicke-Korsakoff syndrome (WKS). Overall, only a small proportion of pregnant women with or without prior thiamine deficiency tend to develop HG, which is linked with potential medicosurgical complications including but not limited to only WE, KS and WKS requiring emergency admission and immediate interventions with regular followup till the end of pregnancy.

2. **AIMS**

This study aimed to review the specific literature on various clinical aspects of HG, WE and KS in women with worsening pregnancies. The
secondary objective was to improve the awareness, emphasis on early diagnosis and immediate multimodal intervention concerning aforesaid sequential syndromes in pregnant women. The significance of this study is to fill the knowledge gap of healthcare providers and users especially in Arabian Gulf countries where the published literature on these sequential syndromes is limited and scanty.

3. METHODS

3.1 Search

Electronic searches (from inception to 2019) of four databases and three open access publishing houses (GoogleScholar, PubMed, ScienceDirect, and OvidSP and MedicalDovepress.com, Hindawi.com and Sciedomian.org) and local literature were conducted using Boolean operators and keywords such as Hyperemesis gravidarum AND thiamine deficiency OR severe nausea AND vomiting OR Wernicke encephalopathy OR Korsakoff syndrome OR central pontine myelinolysis AND interventions OR prevention strategies OR adverse effects OR Case reports OR case series OR observational studies OR randomized clinical trials (RCTs) OR systematic reviews OR meta-analysis for retrieving pertinent articles published in English literature. The searches were modified whenever needed and compatible with databases and publishing houses.

3.2 Search Results

A large number of articles more than twenty one thousand were retrieved. A quick screening by a single author excluded more than sixteen thousand articles that did not focus on HG, thiamine deficiency, WE, KS and WKS and their treatment perspectives. Then, two authors independently reviewed the available data for extracting the most relevant articles. Subsequently, unrelated articles, inaccessible papers because of high price tag, articles cited in systematic reviews and meta-analysis, no abstract available, duplications, and irrelevant information were excluded from this study. The remaining articles were screened further for eligibility and those articles which did not focus on HG, WE, KS and WKS and their etiologies, risk factors, comorbidities, complications and

Fig. 1. Prisma diagram summarizing the flow of search results
treatment interventions were excluded. Thus, the total articles included in this narrative review 144 (Fig. 1).

4. RESULTS

4.1 Thiamine (Vitamin B1)

Thiamine is an essential micronutrient obtained from food and dietary, mineral and vitamin supplements, and its adequate daily supply is associated with good pregnancy outcome [52]. Thiamine deficiency is a nutritional disorder and is determined by multiple factors including medications such as diuretics [53]. Besides maternal characteristics, thiamine deficiency is the core problem in HG and poor outcome of pregnancies. In addition, thiamine infusion rarely causes anaphylactic shock and cardiac arrest [45,53,54]. Thiamine a water soluble vitamin is absorbed from the duodenum and 25-30 mg of thiamine is stored in the body system [55,56]. The recommended daily dose of B1 is 0.4 mg. However, in pregnancy the requirement of thiamine increases up to 1.5 mg/day because of developing fetus and hypermetabolic state of pregnancy. This amounts to about 36% above the non-pregnant level [55-57]. The body has approximately 3-week thiamine stores regardless of body mass index (BMI). Furthermore, thiamine requirement is mostly contingent on metabolic rate and, hence, largely increases during high glucose intake including glucose-containing intravenous fluids. In pregnancy specifically, thiamine stores are rapidly depleted due to excessive vomiting, poor intake of food, and greater metabolic demand [56-59]. Therefore, thiamine deficiency is highly prevalent during pregnancy worldwide especially low and middle-income countries. Simply the body redirects maternal thiamine to the fetus during later third trimester of pregnancy. This results in exacerbation of maternal thiamine deficiency that increases the risk of HG, fetal loss and preeclampsia [58,60-62]. Women with HG require even more thiamine due to their high carbohydrate diet, coexisting electrolyte deficiencies of sodium, potassium and magnesium, limited food varieties, chronic starvation and malnutrition, impaired absorption, and reduced muscle mass for storage [1,44-48,52,56,58,63,64]. In sum, thiamine is mainly found in food and its daily requirement increases in pregnancy attributed to a variety of maternal and fetal factors together with inadequate supply of food varieties across all communities and, hence, thiamine deficiency during deteriorating pregnancies vary across the world [43-48,58,64].

4.2 Thiamine and Central Nervous System (CNS)

Thiamine subserves various biochemical functions in the body and its inadequate supply is reported to perturb the functions of various organs including brain and gut [65]. Normally, thiamine is converted into thiamine pyrophosphate in neuronal and glial cells and serves as an active cofactor for transketolase, pyruvate dehydrogenase, and alpha ketoglutarate enzymes (pentose phosphate pathway). These important enzymes are involved in carbohydrate and lipid metabolism, and production both of amino acids and glucose-derived neurotransmitters [65-70]. In addition, thiamine has an important role in axonal conduction concerning acetylcholinergic (acetylcholine) and serotoninergic (serotonin) neurons/receptors. Various subtypes of 5-hydroxytryptamine (5-HT, serotonin) receptors are reported to correlate with emesis arising from multiple diseases including HG and radiation therapy or chemotherapies [70]. Thus, thiamine deficiency over a period of 2-3 weeks leads to dysfunctions of aforesaid enzymes especially of α-ketoglutarate-dehydrogenase in mitochondrial network. Consequently, oxidative cascade results in diffuse impairment in glucose-energy metabolism with necrotic and apoptotic insult to the specific brain regions with the highest metabolic activity including ocular changes such as intra- and extra-cellular edema, glial cell proliferation, neuronal demyelination, and cellular degeneration [69,71]. Chronic alcohol consumption is also associated with reduced transketolase activity by 35% in the cerebellum attributed to thiamine deficiency [72]. Interestingly, thiamine level in the brain is inversely related to the duration of alcohol use. Additional neurobiochemical findings concerning acute thiamine deficiency include increased cytokines, lactate and edema, extracellular glutamate concentrations, nitric oxide from endothelial cell dysfunction, and deoxyribonucleic acid (DNA) fragmentation in neurons, free radical production and breakdown of the blood-brain barrier [67-72]. The neurological implications of thiamine deficiency are diverse but not limited to WE, KS, WKS and dementia or amnestic syndrome. Central pontine myelinolysis (CPM), akinetic mutism and akinetic-rigid syndrome are due to the correction of thymine-induced sodium deficiency [1,29-31,34,35,73-75].
Most of neurological effects of thiamine deficiency are reversible by thiamine repletion. For example, ocular motor signs concern the reversible lesions in the brainstem affecting the abducens nuclei and eye movement centers in the pons and midbrain [71,75,76]. This is evidenced by the fact that the rapid improvement and nearly complete recovery tend to occur following thiamine intake [75]. Furthermore, thiamine-induced ataxia and stance are caused by damage to the superior vermis in the cerebellum, and the degeneration of the Purkinje cells [76,77]. The effects of thiamine deficiency does not stop here because the vestibular apparatus, diencephalon network linked with medial temporal lobes and amygdala are also affected causing impairment of hearing and persistent amnestic syndrome or WKS; the latter is attributed to irreversible damage [68,75]. In sum, thymine deficiency attributed mainly to HG and chronic alcoholism impacts adversely several molecular processes in the CNS leading to WE, extra-CPM and CPM, KS and WKS.

4.3 Diagnosis and Treatment of Thiamine Deficiency

It is wise to diagnose any health condition prior to starting treatment intervention. Laboratory tests for assessing thiamine deficiency are not straightforward and confirmatory because tests reflect only 0.8%-10% of the body’s thiamine stores and thiamine intake [78]. Nonetheless, all women need laboratory workup concerning vitamin profile because at times well nourished patients are reported to show deficits of thiamine and vitamin C [79]. Furthermore, thiamine testing is not always available at all settings and even its reliability is equivocal [78]. About 50% of patients with WE have normal thiamine levels, which mean an array of other factors cause WE, KS and WKS. Evidently, the end result of multiple factors underpinning thiamine deficiency should be managed proactively by thiamine repletion. In addition, the management of other risk factors underlying thiamine deficiency must be considered earnestly or else health providers do disservice to the patients with NVP, HG, WE and KS [51,54,78]. Without any overemphasis, the treatment of thiamine deficiency in pregnant women should be early, adequate and longterm till the end of pregnancy. Parenteral thiamine is non-toxic and rarely causes anaphylaxis, and, hence, its proactive administration is safe during pregnancy. Thiamine doses, 50mg to 200mg a day (up to 500mg) should be given parenterally for two weeks followed by oral dose of 60 mg/day till the end of delivery [30,55,56,67,68]. In addition, magnesium sulfate 1 to 2 ml of 50% solution IV is also recommended as magnesium is a cofactor in many enzymatic processes in the body [52,64]. Overall replenishing thiamine deficiency in pregnant women protects fetus and mother from developing many adverse outcomes [44-51]. The prognosis is variable and for an acute episode, more than 50% patients show substantial improvement and less than 25% remain impaired [30,55,67,68]. In sum, thiamine deficiency in pregnant population affects several biochemical body functions and needs early diagnosis and prompt treatment for the safety both of the fetus and mother.

4.3.1 Hyperemesis gravidarum

HG is a medical emergency characterized by a constellation of symptoms including nutritional deficiencies and metabolic disturbances in the first trimester (4 to 12 weeks) and caused by hormonal changes and acute or chronic thiamine deficits [2-6]. HG might be predicted by several biopsychosocial markers which are pre-pregnancy body mass index (BMI), adipose tissue, maternal age, leptin, nephatin-1, ghrelin, beta-choriionic gonadotropin (β-hCG), estrogen, progesterone, total (T4) and free thyroxine (fT4), placental growth hormone, prolactin and adreno cortical hormones [80]. A recent study reported the possible role of two genes-growth differentiating factor 15 (GDF 15) and insulin-like growth factor-binding protein 7 (IGFBP7) in HG [81]. The pathophysiology of HG is poorly defined and panoply of risk factors and etiologies are proposed yet this avenue needs further research.

4.3.2 Prevention of HG

Converging evidence suggests that prevention is cost-effective and better than cure. Prior to any intervention, all pregnant women need clinical workup and laboratory tests concerning nutritional deficiencies [79]. Evidently, as vitamin B₁ is rarely associated with anaphylactic shock and cardiac arrest [54] and, hence, physicians should ask pregnant patients with HG about thiamine hypersensitivity. All women at risk for recurrent HG if planning for pregnancy should be prescribed multivitamins and minerals for correcting related deficiencies [52,64,79,81-86]. This proactive approach needs to be used at community level in order to treat as many as women susceptible to develop HG and its adverse consequences [1,31,44-48]. Postconventionally, parenteral vitamins with a
minimum of 5 mg of thiamine and adequate fluids with electrolytes should be given to all pregnant women with nausea, vomiting, multiple pregnancy, and neurological and cardiac symptoms for preventing HG, WE and other complications [8-48,73,82,87-90]. On the contrary, administration of parenteral fluids without vitamins and electrolytes interfere with the prevention of nausea and vomiting in pregnant women leading to various complications including WE, KS, CPM and WKS [78,90]. Besides economic burden, HG also causes depression, post-traumatic stress disorder (PTSD) and anxiety in 80% pregnant women who certainly need psychosocial interventions [91]. In sum, HG is induced by diverse biopsychosocial factors and, hence, requires prevention and treatment for reducing its various complications, economic burden and deaths in pregnant women [21,42,92,93].

4.3.3 Treatment of HG

The pregnant women with HG need urgent admission ("holding function" effect) and prompt treatment. Evidently, up to 90% of pregnant women develop short duration of mild nausea and vomiting of pregnancy (NVP) or emesis gravidaum at 4 to 8 weeks of gestation. NVP occurs commonly in young, uneducated, obese, non-smoker and primigravida women and women with a previous history of nausea and vomiting. A selective literature review reported symptom-based (flank pain, neurological symptoms, diarrhea, and abdominal pain) 34 differential diagnoses including diabetic ketoacidosis of NVP [38]. The severe prolonged symptoms of NVP associated with thiamine deficiency suggest intractable HG [38,94]. The management of grade I (without metabolic changes) and II (with metabolic changes) HG requires multimodal approach including several medications associated with multiple adverse effects (Table 1)[38,95]. Parenteral or enteral rehydration among patients with HG needs careful approach because of development of refeeding syndrome with rhabdomyolysis and diabetes insipidus attributed to fatal shift in electrolytes and fluid balance in malnourished patients [96-99]. Although multiple factors underlie refeeding syndrome, hypophosphataemia is its key determinant [99]. Physicians need to follow the guidelines for treating nausea and vomiting with weight loss (≥5kg) among pregnant women [96,97]. In addition, replenishment of deficiencies concerning vitamin K, thiamine, zinc, selenium, iron, phosphorous, sodium, magnesium and calcium benefits mother and fetus and may preempt additional complications [40-52,64,86,97]. The use of thiamine prior to glucose or post-glucose remains an unresolved issue, though most clinicians prefer to use first thiamine followed by fluids for the prevention of WE [100,101]. Currently, advanced digital HG Care App helps in the management of patients with HG [102]. Besides multimodal interventions [38,95], non-pharmacological integrative therapies such as psychological support, hygienic food, ginger (1gm/day), aromatherapy and medical acupuncture at acupoint P6 (Neiguan) benefit patients with HG [95,103,104]. In a nutshell, prevention and appropriate treatment of HG targeting multiple vitamin and mineral deficiencies, comorbidities and careful fluid balance along with proper nursing care avert the development of potentially serious complications including optic atrophy and visual loss in pregnant women and growing fetus [31,37,40-48,90,105-108] (Table 1).

4.3.4 Wernicke encephalopathy

Carl Wernicke first described WE as polioencephalitis hemorrhagica superioris in 1881. Six years later, Korsakoff explained WE as polyneuritic psychosis [67,68]. Over decades, various sociodemographic risk factors of WE identified among pregnant women included middle age, housewives, low education, poor income, multigravida, multiple gestations, use of contraceptives, history of abortion and hospital admission for NVP and HG [18]. WE is a rare, serious neurological complications of HG (or HG is the cause of WE) and typically identified by a triad of symptoms including ataxia, confusion and nystagmus and gaze palsies [22-29,41,50,71,77,109-114]. Patients with chronic alcoholic WE may not have all signs but any two of the following: dietary deficiencies (nausea, vomiting and thiamine deficiency), oculomotor abnormalities (nystagmus), cerebellar dysfunction (ataxia) and/or either an altered mental state (confusion) or mild memory impairment [115]. Certain patients with non-alcoholic WE may have only nystagmus and vestibulo-ocular reflex abnormalities as the initial presentation [116]. However, 10-47% of pregnant patients with WE lack one or two of these signs specifically when they have gradual or episodic onset of WE progressing to chronic WKS (or caused by HG) and 19% of patients may not have any of these symptoms [73,117,118]. In such scenarios, the diagnosis of WE becomes difficult in non-alcoholic and surgical malnourished patients [33,77,87,119-121]. In the absence of universal diagnostic criteria, WE is often misdiagnosed and
also underdiagnosed when precipitated by acute severe thiamine deficiency, infection and metabolic and endocrine perturbations [25, 77,113,120,122,123]. The case of Charlotte Brontë’s death illustrates the difficulties of making the diagnosis of HG/WE induced by refeeding syndrome [124]. Chronic WE occurs among pregnant patients with mild persistent thiamine deficiency and shares nonspecific symptoms such as headaches, anorexia, irritability and abdominal discomfort with HG. These shared symptoms also posit a diagnostic conundrum, HG versus WE [120,121]. However, persistent or prolonged vomiting, confusion, and unintentional weight loss are red flag signs of chronic WE [77,113,120]. Progressive course of WE is characterized by weakness, nausea, abdominal pain, dysarthria, akinetic-mutism, aphasia, seizures and cardiac failure [34,74,77,113]. Furthermore, mental status changes are virtually universal in WE and exhibited as dizziness, drowsiness, disturbed attention with poor concentration, apathy and cognitive impairment. Gait abnormalities range from weakness to inability to stand and may be difficult to identify in HG patients with vertigo and postural hypotension [77]. The Hyperemesis Impact of Symptoms Questionnaire (HIS) and the Pregnancy-Unique Quantification of Emesis and Nausea (PUQE-12 & 24) scoring indexes, which assess holistically nausea and vomiting in pregnant women may further help in the diagnosis of Wernicke encephalopathy. WE is most commonly associated with chronic alcohol abuse [1,72] but can be caused by other important risk factors [26,66,77,125,126] (Table 2). The suggested thiamine doses are 500 mg per day q.i.d for 2 days, followed by 250 mg per day q.i.d. until the patient tolerates oral thiamine of which dose ranges from 60 to 100 mg daily for at least 3 months [66,109]. Most patients with WE improve rapidly with this regimen, and when there is no response, physicians should look for and manage associated diseases including GIT disorders [8-39,66,94,136-138,139] (Table 2). The prognosis of patients with WE depends on the severity, stage of the disease, hospital admission, and the prompt treatment [110]. Evidently, symptoms of WE take different timeline concerning improvement. A complete resolution of nystagmus is usually observed within 1-2 weeks of thiamine use. However, gait disturbances may persist in up to 60% of patients [140]. In case, Wernicke encephalopathy remains unidentified and untreated may progress to chronic Korsakoff syndrome [141]. WE can also progress to coma and even death, with a mortality rate varying from 20% to 30%, mainly due to pulmonary infection, sepsis and decompensated liver disease [110]. Complete remission of WE is reported only in 29% (54% within 3 months) of the cases and about 34% of patients show residual neurological manifestations [66,109,142]. Wernicke encephalopathy is linked with 50% of fetal deaths attributable to spontaneous abortion, elective abortion, intrauterine death and stroke. However, early treatment with thiamine projects much better prognosis. In sum, WE is a serious complication of and shares many risk factors and etiological pathways with HG (important cause of WE) (Table 2), needs early recognition and treatment with thiamine repletion.
along with other supportive measures for better prognosis and successful outcome of pregnancy.

4.3.6 Korsakoff syndrome

Korsakoff syndrome tends to develop among pregnant women provided co-occurring HG and WE are not managed on time. Although the initial manifestations of KS are variable, this syndrome is characterized by a disproportionate persistent impairment in memory such as anterograde and retrograde amnesia, cognitive dysfunctions in terms of poor attention and concentration, confabulation and constipation. Korsakoff syndrome is a chronic disorder, and caused by chronic depletion of vitamin B₁ attributable to multiple etiological pathways including hyperemesis gravidarum. KS co-occurs with HG, WE as WKS, akinetic mutism and dementia [34,57,75,143]. KS accounts for 10% of dementia and often preceded by an episode of WE but this is not always the case; however, Wernicke-Korsakoff syndrome is attributed to multiple factors including most importantly thiamine deficiency and chronic alcohol abuse [34,75,143]. Chronic KS arising from or caused by HG is an uncommon happening and often misdiagnosed and under-diagnosed condition. A high index of suspicion through the lens of global amnesia and confabulation enables early recognition and, hence, timely treatment with thiamine decreases maternal morbidity and mortality and also improves outcome of pregnancy. Thiamine replenishment and regular followup should continue until patient is asymptomatic. In sum, non-alcoholic KS or WKS caused primarily by chronic thiamine deficit and HG or WE in pregnant patients takes a downhill course with possible permanent neurological disabilities, institutionalization and death when not managed on time by thiamine treatment and supportive measures [75,141,143].

5. DISCUSSION

This review critically described hyperemesis gravidarum, Wernicke encephalopathy, Korsakoff syndrome. Evidently, pregnant women commonly show manifestations of morning sickness (up to 90%) and nausea and vomiting of pregnancy (90%) in early part of gestation, i.e., 3-8 weeks and respond well to prompt intervention with thiamine, life style modification and psychosocial support [2-7]. However, a small proportion of vulnerable pregnant women continue to develop persistent nausea and vomiting with metabolic changes, and ataxia, confusion and oculomotor abnormalities, and gross memory impairment and abdominal disturbances compatible with the diagnostic formulation of hyperemesis gravidarum [2,3,6,17,18,21-28,32], Wernicke encephalopathy [32,66,67,69] and Korsakoff syndrome [34,57,75,141,143], respectively. The relevant literature suggests that women with these sequential syndromes show wide variations in epidemiological trends including risk and predictive factors, etiologies, clinical manifestations, laboratory results, treatment options, and prognosis and outcome mainly attributed to absence of clear definitions, diagnostic criteria, universal treatment guidelines and most importantly methodological differences [38,43,45,51,58,72,80,85,136]. Of note, a wide range of diverse co-occurring systemic diseases and multiple medicosurgical complications contribute further to the complexity of individual syndrome and variable presenta-tions across the board [8-39,34,57,75,143]. In addition, although hormonal changes, increasing metabolic demand, thiamine deficiency, several genes, inflammatory and immunological markers are elucidated in the pathophysiology of HG, WE, KS and WKS yet this avenue needs further research [2-6,51,80,136,144].

This critical review has some limitations including selection and publication biases, because a large number of articles published in closed access journal were not accessible. The accessibility of the relevant studies published in open access journal was also restricted due to high price tag. We also did not attempt to trace unpublished studies by contacting their authors. Despite all these inaccuracies, we synthesize the data from the most influential accessible articles including case reports, observational studies, narrative reviews, updates, systematic reviews and meta-analysis for this critical review. The strength of this study is that it addresses the most important research avenue concerning women’s health in the pregnancies. This review may enhance globally the awareness of sequential syndromes and fill the knowledge gap of professionals providing therapeutic services to pregnant women developing nausea and vomiting of pregnancy, hyperemesis gravidarum, Wernicke’s encephalopathy, Korsakoff syndrome and Wernicke-Korsakoff syndrome. Finally our team suggested a new collective term for these conditions as “B₁-sequential syndromes” because prime cause, clinical picture and treatment interventions are shared by these disorders (Fig. 2).
Table 1. Co-occurring diseases and adverse consequences of HG, adapted from references

<table>
<thead>
<tr>
<th>Diseases</th>
<th>Remarks</th>
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<tbody>
<tr>
<td>Pneumomediastinum and pneumothorax</td>
<td>Serious emergency and needs surgical intervention</td>
</tr>
<tr>
<td>Esophageal rupture or perforation</td>
<td>Serious emergency and needs surgical intervention</td>
</tr>
<tr>
<td>Barogenic rupture of the esophagus</td>
<td>Serious emergency and needs surgical intervention</td>
</tr>
<tr>
<td>Mallory-Weiss syndrome (M-WS)</td>
<td>Control of vomiting to decrease oesophageal pressure</td>
</tr>
<tr>
<td>Renal failure</td>
<td>Serious emergency needing dialysis</td>
</tr>
<tr>
<td>Liver dysfunction</td>
<td>Treatment will depend on severity</td>
</tr>
<tr>
<td>Transient Thyrotoxicosis</td>
<td>Needs intervention</td>
</tr>
<tr>
<td>Visual loss/optic neuropathy</td>
<td>Needs ophthalmic intervention</td>
</tr>
<tr>
<td>Hamman and Boerhaave syndrome</td>
<td>Needs intervention</td>
</tr>
<tr>
<td>Thiamine deficiency</td>
<td>Repletion needed</td>
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<tr>
<td>Iatrogenic WE</td>
<td>Fluid load correction</td>
</tr>
<tr>
<td>Korsakoff syndrome and WKS</td>
<td>Thiamine plus care of other factors</td>
</tr>
<tr>
<td>Extraintestinal and CPM</td>
<td>Correction of electrolytes and fluid</td>
</tr>
<tr>
<td>Akinetic mutism,</td>
<td>Thiamine and supportive measures</td>
</tr>
<tr>
<td>Akinetic-rigid syndrome</td>
<td>Thiamine and other treatment</td>
</tr>
<tr>
<td>Locked-in syndrome</td>
<td>Thiamine and other treatment</td>
</tr>
<tr>
<td>Chronic alcohol abuse</td>
<td>Needs intervention with thiamine</td>
</tr>
<tr>
<td>Infection, metabolic and endocrine perturbations</td>
<td>Acute thiamine deficiency needs multimodal interventions</td>
</tr>
<tr>
<td>Starvation and nutritional deficiencies</td>
<td>Correction of deficiencies through feeding</td>
</tr>
<tr>
<td>Electrolyte imbalance (Low K &amp; Na)</td>
<td>Correction of sodium &amp; potassium after thiamine use</td>
</tr>
<tr>
<td>Acute pancreatitis</td>
<td>Treatment of infection/inflammation and fluid balance</td>
</tr>
<tr>
<td>GIT infection (HP)</td>
<td>Needs antibiotic (erythromycin) intervention</td>
</tr>
<tr>
<td>Low mineral (phosphorous)</td>
<td>Needs urgent correction</td>
</tr>
<tr>
<td>Muscle weakness and tetany</td>
<td>Dietary supplements and electrolyte replenishment</td>
</tr>
<tr>
<td>Acidosis and alkalosis (due to malnutrition &amp; vomiting)</td>
<td>Needs correction by fluids</td>
</tr>
<tr>
<td>ECG abnormalities</td>
<td>Intervene if needed</td>
</tr>
<tr>
<td>Fetal growth retardation and death</td>
<td>Monitoring and careful disposal</td>
</tr>
<tr>
<td>Psychological impact on quality of life, occupation, body image, relation with spouse, absenteeism, cost burden, job loss, depression, anxiety and PTSD</td>
<td>Counselling, psychotherapy, hypnotherapy, behavioral therapy including CBT and specific medications</td>
</tr>
<tr>
<td>Retinal hemorrhage</td>
<td>Need ophthalmic intervention</td>
</tr>
<tr>
<td>Epistaxis (due to low Vitamin K)</td>
<td>Needs vitamin K</td>
</tr>
<tr>
<td>Vasospasm of cerebral arteries</td>
<td>Needs urgent treatment of HG</td>
</tr>
<tr>
<td>Preeclampsia</td>
<td>Immediate intervention</td>
</tr>
<tr>
<td>Peripheral neuropathy</td>
<td>Vitamin supplements</td>
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<tr>
<td>Coagulopathy</td>
<td>Medications or replacement therapy directed toward absent clotting factors</td>
</tr>
<tr>
<td>Rhabdomyolysis and diabetic insipidus</td>
<td>Fluid resuscitation and prevention of end-organ complications such as kidney failure by dialysis</td>
</tr>
</tbody>
</table>

Multimodal Approach: Antiemetics (dimenhydrinate 25-50 mg every 6-8 h; meclizine 25-100 mg 2-4× per day orally/1× per day rectally; dimenhydrinate: 50 mg 3-4× per day orally/62 mg 2× per day IV/1-3× per day rectal route; promethazine: 12.5-25 mg 6-8 hourly orally, IM or IV and doxylamine (oral 25 mg at night and 12.5 mg in the morning along with 10 mg to 80 mg pyridoxine); phenothiazines (Prochlorperazine 25 mg per day or 2×per day rectally; Chlorpromazine: 5-10 mg 6 hourly orally, IM or IV; 25 mg per rectally; metoclopramide, 5-10 mg 8 hourly orally, IV or IM (maximum 5 days’ duration); steroids (hydrocortisone 100 mg twice daily, then 50mg/day and then tappers lowly); benzodiazepines (diazepam 40-160 mgIV, or midazolam 1 to 2mg IV slowly over 10 minutes); 5HT3 antagonist (Ondansetron: 4-8 mg 6-8 hourly PO; 8 mg over 15 minutes 12 hourly IV); thiamine (50 to 200 mg/day parenteral); pyridoxine (Vit.6, 10-25 mg tid and maximum dose 200 mg/day); enteral and parenteral fluids (3L/day) and carbohydrate and amino acid (about 8400 to 10,500 kJ/d) with complete withdrawal of oral feeding.
Table 2. Etiological and risk factors of HG (and Wernicke’s encephalopathy)*, adapted from references

<table>
<thead>
<tr>
<th>Etiological and risk factors</th>
<th>Adapted from references</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excessive, prolonged vomiting in pregnancy including past history</td>
<td>Severe infection and malnutrition due to child abuse, fasting and extreme dieting and hunger strike</td>
</tr>
<tr>
<td>Current and previous molar pregnancy</td>
<td>Pre-existing diabetes and diabetic ketoacidosis</td>
</tr>
<tr>
<td>Eating disorders; anorexia nervosa, bulimia nervosa and binge eating</td>
<td>Malabsorption</td>
</tr>
<tr>
<td>Chronic kidney disease and pyelonephritis</td>
<td>Glucose/dextrose infusion prior to thiamine administration</td>
</tr>
<tr>
<td>Haemodialysis or peritoneal dialysis</td>
<td>Acute or chronic thiamine deficiency</td>
</tr>
<tr>
<td>Chemotherapy</td>
<td>Inadequate thiamine intake</td>
</tr>
<tr>
<td>Gestational thyrotoxicosis</td>
<td>Antibiotics</td>
</tr>
<tr>
<td>Prolonged parenteral feeding and TPN, prior to thiamine replenishment</td>
<td>Diuretics</td>
</tr>
<tr>
<td>Multiple gestation</td>
<td>Genetic factor; placentation, appetite, and cachexia genes</td>
</tr>
<tr>
<td></td>
<td>GDF15 and IGFBP7 are linked to hyperemesis gravidarum (HG) and are upregulated in HG patients.</td>
</tr>
<tr>
<td>Anemia</td>
<td>Chronic alcohol abuse and non-smokers, smokers show low levels of estrogen and HCG.</td>
</tr>
<tr>
<td>Gastroplasty</td>
<td>Malignant disorders and paraneoplastic disease (signs and symptoms as a consequence of cancers)</td>
</tr>
<tr>
<td>GIT disorders, pyloric stenosis, lower esophageal sphincterdis order, cancer, infection (HP), gastrectomies, delayed gastric emptying and intestinal transit times</td>
<td>Acquired immunodeficiency syndrome</td>
</tr>
<tr>
<td>Iron substitution</td>
<td>Organ transplantation (liver), cholecystitis and hepatitis</td>
</tr>
<tr>
<td>Thyroid dysfunction (transient hyperthyroidism with↓ level of TSH persists only first 18 weeks of pregnancy)</td>
<td>Hormones—estrogen (↑) and beta-HCG, IgG and IgM &amp; prostaglandin E2, progesterone (↑ or ↓), causing gastric dysrhythmias - tachygastria, or bradygastria, ACTH, cortisol, growth hormone, 5HT and prolactin.</td>
</tr>
<tr>
<td>Unwanted pregnancy,</td>
<td>Psychosomatic model; Psychiatric disorders especially depression, conversion disorder; sequel of HG rather than vice-versa</td>
</tr>
<tr>
<td>Female sex of offspring,</td>
<td>High saturated fat diet</td>
</tr>
<tr>
<td>Immunological factors (high levels of immune globulins, C3, C4, and lymphocyte Counts)and autoimmune diseases</td>
<td>Nulliparous woman</td>
</tr>
<tr>
<td>Obesity</td>
<td>H/O Migration</td>
</tr>
<tr>
<td>Drug intoxication</td>
<td>Food poisoning</td>
</tr>
<tr>
<td>Residence in a nursing home</td>
<td>Habituallty restricted diet in the elderly living alone</td>
</tr>
</tbody>
</table>

*other comorbid conditions are acute disseminated encephalomyelitis, pituitary apoplexy (bleeding into pituitary gland and or impaired blood supply), deep cerebral vein thrombosis, pancreatitis, multiple sclerosis and Miller Fisher syndrome

HP=Helicobacter pylori in 95% women with intractable HG and higher IgG against HP; HCG=human chorionic gonadotrophin; ACTH=adrenocorticotropic hormone; TSH=thyroid stimulating hormone.
6. CONCLUSION

Pregnancy is a normal physiological condition with increasing metabolic demand and hormonal changes and progresses to normal delivery ending at 9-month with live infant. However, certain vulnerable pregnant women develop morning sickness, NVP, HG, WE, KS and WKS. The prime cause underlying these sequential syndromes is thiamine deficiency but a variety of other etiological and risk factors are evoked to explain the pathophysiology of individual syndrome; however, till now these conditions remain poorly understood. Each pregnant patient with HG or WE or KS or WKS is unique and needs comprehensive clinical diagnostic workup to exclude various systemic diseases using various means including brain MRI. Most pregnant patients with NVP, HG, WE, KS and WKS tend to improve with multimodal treatment that needs to be continued till the successful delivery; however, a variable number of patients with WE, KS and WKS continue to show residual features. Future studies should focus on strategies as how to globally define these sequential syndromes, develop universally accepted diagnostic criteria for early identification and optimal management, explore inflammatory and genetic markers and improve robustly the maternal morbidity and fetal outcome in the pregnancies.

CONSENT

It is not applicable.

ETHICAL APPROVAL

It is not applicable.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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