



Guidelines for the Management of Hyperemesis Gravidarum

A Clinical Guideline

For Use in:	Obstetrics and Gynaecology		
By:	Clinical staff		
For:	Management of Hyperemesis Gravidarum		
Division responsible for document:	Women and Children's Services		
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If Yes - does the strategy/policy deviate from the recommendations of NICE? If so why?	No		

Version and Document Control:

Version Number	Date of Update	Change Description	Author	
4	02/06/2021	Amended following an incident Prochlorperazine section amended from IM/IV to IM only	Neeraja Kuruba	
5	18/01/2022	Changes as per the new RCOG guideline, changes to key people	Claire Wells, Kelly French, Michelle Drolet, Bryony Tomlinson	

This is a Controlled Document

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Background

Nausea and vomiting in pregnancy (NVP) affects up to 80% of pregnant women and is one of the most common indications for hospital admission among pregnant women. It typically starts between the 4-7th weeks of pregnancy, peaks around 9th week, and resolves by the 20th week in 90% of women. Rule out other causes if first presentation is after 10 completed weeks of pregnancy.

This condition is known as hyperemesis gravidarum which can be defined as intractable vomiting associated with loss of more than 5% of pre pregnancy weight, dehydration, electrolyte disturbances, or need for hospital admission. There is a high risk of recurrence in subsequent pregnancies. Consider diabetic ketoacidosis as an alternative diagnosis in a ketotic woman with diabetes.

Risk factors and associations:

- First pregnancy
- Multiple pregnancy
- History of severe nausea and vomiting in previous pregnancies, motion sickness, or nausea with oral contraceptive use
- Gestational Trophoblastic disease (GTD), including molar pregnancy
- History of migraines
- History of first degree relative with NVP
- Obesity
- Stress
- Being seropositive for Helicobacter pylori

The condition spontaneously resolves in the vast majority of patients and complications are rare.

Complications

- Weight loss
- Electrolyte imbalance
- Abnormal LFTs
- Abnormal TFTs
- Central pontine myelinolysis (CPM)
- Wernicke's encephalopathy
- Other vitamin deficiencies (± megaloblastic anaemia), such as B12 or B6
- Venous thromboembolism (VTE)
- Adverse pregnancy outcomes including low birth weight, and increased risk of preterm delivery
- Adverse effect on quality of life and mental health
- Mechanical complications including Mallory-Weiss tears, retinal haemorrhage

Management

HARP Criteria

Patients that fulfil the criteria should be commenced on the integrated care pathway (ICP) pathway.

- Ketones of 2 or more
- No complications ie: weight loss, deranged bloods
- Consideration of past medical history and comorbidities
- Patient compliance

<u>History</u>

Quantify severity

To exclude other causes: abdominal pain, urinary symptoms, infection, drug history

Examination

Undertake an examination as per the ICP pathway. Includes basic observations, weight, abdominal examination, signs of dehydration, urine output, and other examination as guided by history.

Investigations:

On admission all patients require:

- Urine dipstick to quantify ketonuria
- MSU to exclude UTI if any positive findings on urine dip
- U&Es to identify electrolyte imbalance
- FBC and CRP to exclude infection
- Random blood glucose
- For diabetic patients, checking capillary blood glucose regularly is mandatory

On second admission:

Arrange ultrasound scan via EPAU to confirm viable intrauterine pregnancy, and exclude multiple pregnancy or trophoblastic disease.

Repeated attendances:

In addition to bloods as per first admission which should be checked on each attendance, periodically check TFTs, LFTs, blood group, magnesium, and amylase. Suggested frequency at least every second attendance, or more frequently if abnormal. TFTs may be checked less frequently such as once in first trimester and once in second trimester unless any abnormality noted.

Treatment

Inpatient management should be considered if there is at least one of the following:

- a. continued nausea and vomiting and inability to keep down oral antiemetic's
- b. continued nausea and vomiting associated with ketonuria and/or weight loss (greater than 5% of body weight), despite oral antiemetic's
- c. Confirmed or suspected comorbidity (such as urinary tract infection and inability to tolerate oral antibiotics).
- d. Weigh patient on admission, then twice weekly.

Intravenous (IV) infusions

- Insert non-ported cannula for IV access
- The rate of rehydration depends on the severity of NVP but usually aggressive rehydration with 1 litre of 0.9% saline with 20mmol potassium chloride (KCI) over 2 hours is often appropriate
- 0.9% saline with additional potassium chloride in each bag, guided by daily monitoring of electrolytes is the most appropriate IV hydration regimen as per RCOG guidance
- If hypokalaemic, 20-40mmol of KCl in 0.9% saline (note maximum infusion rate 10mmol/hr of KCl)
- Dextrose infusions are **not** appropriate unless the serum sodium levels are normal and thiamine has been administered
- Avoid glucose as it can precipitate Wernicke's encephalopathy.

Antiemetics

- There are safety and efficacy data for first line antiemetics such as H1 receptor antagonists and they should be prescribed first when required. Although generally safe in the first trimester, they are currently not licensed for use in pregnancy in the UK.
- Combination of drugs from different classes should be used in women who do not respond to a single antiemetic
- Use all antiemetics regularly rather than PRN
- The parenteral or rectal route may be necessary and more effective than the oral regimen for women with severe or persistent NVP

Recommended antiemetic therapies and dosages:

Drug	Dose	Major side-effects	Class	
First Line		-	-	
Cyclizine	50mg PO/IM/IV TDS	Drowsiness, dizziness	H1 receptor antagonist	
Prochlorperazine	5-10mg 6-8 hourly PO (also available as oral solution), 12.5mg 8 hourly IM, 3-6mg 12 hourly buccal	Hypotension, extrapyramidal symptoms (tardive dyskinesia, dystonia)	Antipsychotic phenothiazines	
Promethazine (Phenergen)	12.5-25mg 4-8 hourly PO /Deep IM	Drowsiness, sedation	H1 receptor antagonist	
Chlorpromazine 10-25mg 4-6 hourly PO/ Deep IM Sedation, symptoms		hypotension, extrapyramidal	Antipsychotic phenothiazines	
Second Line				
Metoclopramide	5-10mg 8 hourly PO/IV/IM (also available as oral solution)	Extrapyramidal symptoms (torticollis, oculogyric crisis)	D2 receptor antagonist	
Domperidone	10mg 8 hourly PO, 30-60mg 8 hourly PR	Minimal	D2 receptor antagonist	

Second line if >13 weeks				
Ondansetron	4-8mg 8 hourly PO/IV	Headache, GI upset, Should be given >13 weeks only due to fetal risk of cleft palate and renal abnormalities	5-HT3 receptor antagonist	
Third Line	•	•		
Corticosteroids Hydrocortisone 100mg BD IV, convert to prednisolone 40- 50mg OD PO and taper dose to lowest level which still controls symptoms		Patients should be given clear advice about self- management of tapering, and a steroid alert card		

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When all other medical therapies have failed, enteral or parenteral treatment should be considered with a multidisciplinary approach. In refractory cases artificial nutritional support should be considered. Enteral (nasogastric or nasojejunal) or parenteral feeding can be considered. The MUST screening tool may be useful. Enteral feeding is contraindicated in women with acute vomiting due to the risk of aspiration ensure correct procedure

<u>Thiamine</u>

- Should be given to all women attending with prolonged vomiting, to prevent vitamin deficiency and Wernicke's encephalopathy.
- Thiamine 50mg oral once daily should be given to all patients treated according to the HARP for a 10 day course.

<u>Iron</u>

• Consider avoiding iron-containing preparations if these exacerbate symptoms

Anti-GORD measures

• Proton pump inhibitors (e.g. omeprazole) or H2 receptor antagonists may be used for women developing gastro-oesophageal reflux disease (GORD). Both are considered to be safe in pregnancy.

Venous Thromboembolism (VTE)

• Perform VTE risk assessment as per the VTE chart in the HARP integrated care pathway and treat accordingly. Appendix A

Patient education

- Reassurance
- Rest
- Dietary and lifestyle advice
- Patient information leaflet e.g.

https://www.rcog.org.uk/globalassets/documents/patients/patientinformation-leaflets/pregnancy/pi-pregnancy-sickness.pdf https://www.pregnancysicknesssupport.org.uk/resources/printableleaflets/

- Signpost to additional support and information e.g.
 Pregnancy Sickness Support: <u>www.pregnancysicknesssupport.org.uk</u>
- NHS Choices

Impression:

Based on this assessment the patient may either be suitable or not suitable but should be considered for the HARP, as per inclusion and exclusion criteria above. If the patient is suitable for HARP, follow the rest of this guideline and commence with Integrated Care Pathway (ICP). Appendix B

Patients that fulfil the HARP criteria

<u>HARP</u>

- Commence integrated care pathway (ICP) for HARP
- If inpatient arrange readmission for the following day and discharge with venflon in situ
- If emergency admission arrange immediate commencement of HARP and cannulate
- Follow appropriate day as per ICP

Monitoring

- Observations 4 hourly unless otherwise indicated by the NEWS score
- Input / output chart
- Weight recorded each admission

Discharge Criteria

- Patient understands and agrees to comply with discharge instructions including cannula care if relevant
- Patient has a supply of oral antiemetic's to take home
- Satisfactory observations/ NEWS score
- Tolerating fluids and some food
- Passing adequate urine (0.5ml/kg/hr minimum)
- Make arrangements for follow up as below
- Ensure discharge criteria met for discharge home
- Ensure discharge advice given and provide open access information.
- Open access will continue until the patient reaches 22 weeks gestation at which point care must be transferred to the obstetric team
- Telephone number for Cley ward
- Give the patient the information leaflet on NVP
- Ensure that the patient has received the HARP cannula care leaflet

• Consider giving a supply of urinalysis strips for home ketone testing if felt to be appropriate in individual cases

Criteria for Mandatory Doctor Review:

- Persistent ketonuria on 4th day
- Abnormal NEWS
- Failure to control symptoms
- Deranged biochemistry
- Complications/ development of new problems
- At nurse's request for advice or review
- Where the nurse deems the patient to benefit from doctor review, the nurse will request the SHO/ Registrar/ Consultant to take over the care of the patient and document this in writing.
- After day 4 for admission or review of restarting HARP

Clinical audit standards / audit standards

To ensure that this protocol is compliant with the standards set out a random sample of 10 ICPs will be audited annually to ensure they are completed accurately. The audit results will be sent to the gynaecology matron who will review the audit standards and make recommendations for further actions

Summary of development and consultation process undertaken before registration and dissemination

During the development process the protocol had been circulated between members of the gynaecology department. This included consultants, junior doctors, senior nurses, nurse sonographers and the early pregnancy assessment unit.

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Appendix A

Thromboprophylaxis risk assessment

Thromboprophylaxis risk assessment							
Lower risk (score 1 each)			Higher risk (score 3)				
Less than 3 of the following r	sk factors		Any of the following risk factors		factors		
Age >35 years				Personal history of VTE			
Varicose veins				Med	ical comorbidity e.g.		
Parity ≥3				infla	inflammatory conditions,		
BMI ≥30 (score 2 if BI	VI ≥40)	1		hear	heart/lung disease, SLE, I		
Smoker			type 1 diabetes with		vith		
Current systemic infe	ction	1		nephropathy, nephrotic		ohrotic	
				sync	Irome, cance	er, sickle cell	
				_	ase, IV drug		
Multiple pregnancy/as					Hyperemesis / ovarian		
reproductive techniqu	е				erstimulation	syndrome	
					recovered		
Dehydration					Any Thrombophilia e.g.		
Immobility/journey >4					antiphospholipid syndrome,		
	Family history of unprovoked				Factor V Leiden, Protein C or		
	or <u>oestrogen related</u> VTE in				deficiency, antithrombin		
	first degree family member				deficiency, Prothrombin gene		
	where thrombophilia testing not				mutation		
	performed or results not			Any surgical procedure in			
	available, if not done perform		pregnancy (except SMM or TOP)		pt Sivilvi or		
	antenatal Thrombophillia screen profile bloods. (Blood				Any 3 or more of the 'lower		
profile available on W							
Risk assessment results		risk' factors					
					Lline riel: (T		
Low risk	Intermediate risk			High risk (TRA 4+)			
(TRA <2) No VTE	(TRA 2-3)						
	Prescribes TEDS if			Prescribe LMWH from first trimester and			
prophylaxis required	inpatient, and LMWH for 10 days			continue until 6 weeks			
required			ays		postnatally		
Completed by:					positiatally		
(Print name, signature,					Date:		
designation)					2400		

Appendix B

Integrated Care Pathway to be updated

Appendix C

Patient Information Leaflet for Hyperemesis Ambulatory Rehydration Program (HARP) <u>Trust Docs Id: 18658</u>