Routine Antenatal Assessment in the Absence of Pregnancy Complications

A woman’s health during her pregnancy is critical to the outcome of the pregnancy and may have a lifelong impact on her baby’s health.

FIRST ANTENATAL VISIT IN PREGNANCY

All women should be advised to attend in early pregnancy with a view to:
1. confirming pregnancy and establishing an estimated date of confinement (albeit that may alter after subsequent ultrasound examinations);
2. a comprehensive clinical assessment in order to determine any clinical conditions that may be of relevance to the pregnancy;
3. detailed assessment of any particular conditions or circumstances of relevance and optimising management for pregnancy;
4. obtaining general advice regarding common issues of concern in early pregnancy and management of the pregnancy.

Clinical assessment
As always, of greatest importance is a careful medical history and thorough clinical examination.

The following investigations are recommended (in the absence of specific complications):

Full blood examination
Particular note should be taken of the Mean Corpuscular Volume as a potential indicator of an underlying Haemoglobinopathy.

Blood group and antibody screen
Where the blood group has already been performed it does not need to be repeated. However, the antibody screen should be repeated at the beginning of each pregnancy.

Rubella antibody status
All women should have their rubella antibody titre measured for each pregnancy. Although the past antibodies titres from a previous pregnancy screens may have been used to exclude a further antenatal test, there is evidence that levels may decline, particularly following immunization as compared to natural infection. This is particularly so given the low level of wild virus circulating in the community to boost women whose levels may fall below that of protection.

Syphilis serology
Syphilis testing should be performed by screening with a specific treponema pallidum assay for example Treponema pallidum haemaglutination assay (TPHA) or the Treponema pallidum particle assay (TPPT). The non-specific Treponema pallidum assays, such as rapid plasma regain (RPR) test, although cheaper, are less likely to pick up latent infection.
Midstream urine
Examination by culture, e.g. dip slide.

HIV
Before instituting screening for any viral infection in pregnancy, it is imperative that the woman is provided with appropriate counselling as to the limitations of screening for viral infections in pregnancy and the implications of both positive and negative findings. All pregnant women should be recommended to have HIV screening at the first antenatal visit.

Hepatitis B serology
All pregnant women should be recommended to have Hepatitis B screening in pregnancy.

Hepatitis C serology
All pregnant women should be recommended to have Hepatitis C screening in pregnancy. However it is acknowledged that this is a contentious area of practice.

Varicella
Consideration should be given to checking varicella antibodies at the first visit where there is no history or uncertain history of previous illness.

Cervical cytology
A cervical (Pap) smear should be recommended at the first antenatal visit if this would fall due during the pregnancy, according to cervical screening guidelines. There is no evidence to suggest that a Pap smear in pregnancy is harmful.

OTHER TESTS THAT MAY BE CONSIDERED

Screening for haemoglobinopathies
Each unit should have a defined policy for screening for haemoglobinopathies, taking into account the ethnic mix of patients screened. As a minimum, all women should be screened with MCV and MCHC. Haemoglobin electrophoresis and iron studies should be performed in the event of thresholds not being reached. Consideration should also be given to the further screening of patients with DNA analysis for alpha-thalassaemia. Testing of normal-MCV women for haemoglobinopathies may be considered if they are members of high-risk groups.

Vitamin D
Pregnant women at risk for vitamin D deficiency should be tested in early pregnancy OR provided with vitamin D supplementation.

CMV
Screening for CMV infection in pregnancy is currently not recommended as a routine (see consensus statement on CMV in pregnancy).

General advice
All women in early pregnancy should be informed with respect to:
1. potential teratogens (medications, alcohol, X-rays etc);
2. vitamin and mineral supplementation (see college statement);
3. model of care, expected visit frequency, place of booking for confinement, expected costs for both pregnancy and confinement.

SUBSEQUENT VISITS DURING THE ANTENATAL CARE
All women should be advised to attend in early pregnancy with a view to:
1. early diagnosis of pregnancy complications;
2. utilising the principles of preventative medicine to minimise the risk of problems in pregnancy, labour and the puerperium;
Clinical assessment
All women should have a directed clinical assessment at each antenatal visit, with a focus on general well-being and early diagnosis of pregnancy complications. Investigations recommended are:

Obstetric ultrasound scan
All women should be offered an obstetric ultrasound before 20 weeks' gestation. This will include an ultrasound for fetal morphology and placental localization usually at 18-20 weeks gestation. Other scans may be indicated depending on individual circumstances and to assess/confirm dates.

Screening for Down syndrome
Refer to Antenatal Screening for Down syndrome and other fetal aneuploidy (C-Obs 4), see link below.

Gestational diabetes
Screening for Gestational Diabetes Mellitus is recommended in all pregnant women. See the original (1998) ADIPS guidelines.

Group B Streptococcal Disease (GBS)
Refer to Swabbing for Group B Streptococcus (C-Obs 19), see link below.

Blood group antibody testing
Refer to Guidelines for the use of Rh-D immunoglobulin (anti-D) in obstetrics in Australia (C-Obs 6), see link below. Further screening is recommended for Rh negative women at approximately 28 weeks gestation. Screening of Rh positive women at 28 weeks gestation is at the discretion of the clinician/managing health service.

Iron deficiency
The haemoglobin level and platelet count should be repeated at 28 weeks gestation. If anaemia is detected, further investigation is warranted.

Cytomegalovirus/Toxoplasmosis
Selective testing for cytomegalovirus and toxoplasmosis is recommended only for those women at a substantially increased risk of acquiring an infection. Ideally such patients should be tested prior to pregnancy.

Syphilis
Syphilis screening should be repeated at 28 weeks in high-risk populations.

LATE PREGNANCY TESTS OF FETAL WELL-BEING

Late pregnancy tests for assessment of feto-placental function should be performed when indicated on clinical grounds - either through a suspicion of placental insufficiency, a predisposing factor for placental insufficiency or through an inability to clinically ascertain fetal growth (e.g. obesity). Tests of fetal wellbeing should be considered after 41 weeks' gestation. Detailed and frequent assessment of fetal wellbeing, including an assessment of liquor volume, is mandatory in pregnancies at or beyond 42 weeks gestation.

Chlamydia
Selective testing for Chlamydia should be considered for those who may be at increased risk (e.g. less than 25 years).
Links to other related College Statements

(C-Obs 03a) Pre-pregnancy Counselling

(C-Obs 04) Antenatal screening for Down Syndrome and other fetal aneuploidy

(C-Obs 06) Guidelines for the use of RhD immunoglobulin (anti-D) in obstetrics in Australia

(C-Obs 07) Diagnosis and management of gestational diabetes

(C-Obs 19) Swabbing for Group B Streptococcus

(C-Obs 25) Vitamins and Mineral Supplements in Pregnancy

(C-Obs 45) Influenza Vaccination during Pregnancy

(C-Obs 46) Testing of serum TSH levels in pregnant women

(C-Gen 02) Guidelines for consent and the provision of information regarding proposed treatment

(C-Gen 03) Hepatitis B

(C-Gen 04) Hepatitis C

(C-Gen 15) Evidence-based Medicine, Obstetrics and Gynaecology

Patient Resources

RANZCOG patient information pamphlet:
Antenatal care and routine tests during pregnancy - a guide for women (July 2002).

Other suggested reading


Disclaimer

This College Statement is intended to provide general advice to Practitioners. The statement should never be relied on as a substitute for proper assessment with respect to the particular circumstances of each case and the needs of each patient.

The statement has been prepared having regard to general circumstances. It is the responsibility of each Practitioner to have regard to the particular circumstances of each case, and the application of this statement in each case. In particular, clinical management must always be responsive to the needs of the individual patient and the particular circumstances of each case.

This College statement has been prepared having regard to the information available at the time of its preparation, and each Practitioner must have regard to relevant information, research or material which may have been published or become available subsequently.

Whilst the College endeavours to ensure that College statements are accurate and current at the time of their preparation, it takes no responsibility for matters arising from changed circumstances or information or material that may have become available after the date of the statements.