Antiphospholipid syndrome and pregnancy loss
We have written this leaflet for women and their partners, who have had recurrent miscarriages and have also been diagnosed with antiphospholipid syndrome.

But you might be reading it because you are about to have investigations following recurrent miscarriage, or simply because you want to know more about this cause of miscarriage.

Whatever your circumstances, we hope that you find this leaflet helpful.

What is antiphospholipid syndrome?
The first thing to know is that antiphospholipid syndrome is also called

- Antiphospholipid antibody syndrome or
- Hughes’ syndrome or
- sticky blood syndrome or
- APS

In this leaflet we will usually call it APS.

In general, a syndrome means a combination of signs, symptoms and/or test results. Antiphospholipid syndrome (APS) is the association between three things:

- antiphospholipid antibodies (often written as aPL),
- blood clots in the veins, arteries or small blood vessels (doctors call these thromboses) and/or
- pregnancy problems, including miscarriage

How is it diagnosed?
You will have blood tests to check for antiphospholipid antibodies (aPL). They will actually test the blood for three types of antibody:

- lupus anticoagulant (LA)
- anticardiolipin antibody (aCL) and
- anti beta-2 glycoptein I antibodies.

Don’t be confused by the first of these, “lupus anticoagulant”, which is a rather misleading term.

It doesn’t mean you have the chronic condition called lupus. And it doesn’t mean you have an anticoagulant (blood thinner) in your blood.

The tests should be done twice, with an interval of at least twelve weeks between each test.

This is because aPL can occur just for a short time because of infection (e.g. chickenpox) or some medicines (e.g. phenytoin).

Another reason for repeating the test is that antibody levels vary over time.
It was a relief that there was finally a name for the problem, some explanation for my miscarriages.

What’s more, different laboratories differ in the way they carry out the tests. That means you might test positive for aPL in one hospital and negative in another. This can be confusing for you and your doctors.

You will be diagnosed with APS if you have:

- two positive tests for aPL, taken around 12 weeks apart and one or more of the following:
  - a history of recurrent first trimester miscarriage (three miscarriages in a row), with no other cause being found;
  - previous pre-eclampsia and intrauterine growth restriction, resulting in a baby being born before 34 weeks
  - a second or third trimester pregnancy loss or stillbirth
  - a previous thrombosis (blood clot)

How does APS cause pregnancy problems?

The aPL antibodies can cause different problems at different stages of the pregnancy in different women.

In some women with aPL, in the first 13 weeks, the antibodies can prevent the pregnancy from embedding properly in the womb, increasing the chances of an early miscarriage.

In the second and third trimesters of pregnancy (from 14 weeks gestation until birth), aPL in some women can cause blood clots in the placenta. This can lead to poor blood and oxygen supplies to the baby, causing poor growth, pre-eclampsia or even stillbirth.

Some women with aPL will have no problems in pregnancy at all.

Researchers are still trying to find out more about the link between APS and pregnancy loss.
Can APS cause any particular symptoms or problems?

If you are diagnosed with APS, you may well have no other symptoms or conditions. Some people with APS, however, have one or more of the following:

**Skin**  Approximately one fifth of people with APS have a blotchy, mottled appearance to their skin

**A tendency to bruise easily**  This is unusual. Occasionally women with APS have a low platelet count (platelets are cells in the blood that help the blood to clot)

**Blood clots in the vein**  The most common blood clot in the vein is a deep vein thrombosis (DVT) of the leg.

**Joint pain**  There may be a history of joint pain or swelling in the joints (arthritis).

**Neurological problems**  Some people with APS have migraine-like symptoms or brief loss of vision or balance, or even stroke.

**Abdominal or heart disease**  Rarely, APS can cause liver, heart or kidney problems, due to blood clots. If you are diagnosed with APS, you may be referred for tests such as an echocardiogram of your heart, to check that you are healthy.

How is APS treated?

Treatment of APS depends on your history, test results and whether or not you are planning to become pregnant. In this leaflet, we'll assume that you have had repeated miscarriages, have been diagnosed with APS, but have not had a thrombosis\(^1\) – and that you are thinking about trying again.

Your treatment will be aimed at preventing blood clots. You will almost certainly be treated with low doses of aspirin (usually 75 mg a day) starting early in pregnancy or perhaps even before you conceive. You may also be advised to have injections with another blood-thinning drug, called heparin. There are many different makes of heparin.

Your doctor will advise you when this treatment should be stopped – and whether you need any further treatment or follow-up after your pregnancy.

\[^1\] If APS patients have a thrombosis (blood clot), they are usually treated with warfarin.
Your feelings

Miscarriage can be a very unhappy and frightening event in any circumstances. But if you have experienced recurrent miscarriage, you may have found it much harder to bear than your first or even second loss. You may wonder whether you’ll ever have a baby.

In addition, the investigation process itself can be stressful. You may have had to wait to be referred for tests, then waited for your appointment – and then had to wait for the tests to be repeated over several weeks.

What does having APS mean for the future?

You might be relieved that you’ve been told you have APS. Finally there’s an explanation for your miscarriages and a treatment plan for the future. There is good reason to feel positive – having treatment for your APS makes it much more likely that you’ll have a healthy pregnancy.

On the other hand, you may find it difficult to feel positive.

You have been disappointed so many times before that it can be hard to begin another pregnancy with any hope that it might succeed.

And trying not to be too hopeful may feel like the best way of protecting yourself in case things go wrong again.

Many couples go through a mixture of hope and anxiety – up one minute and down the next – and you might too.

Once you are pregnant, you may find it helps to have regular check-ups to make sure everything is developing as it should. And if you have someone you can call on when you are feeling anxious, this can help too. (See page 7 for some suggestions.)
What if things go wrong?

Sadly, even with expert care and treatment, you might miscarry again, or lose the baby at a different stage of pregnancy.

This might be because the treatment hasn’t prevented your APS from harming the pregnancy. In this situation, your doctor may try a different treatment next time.

Or it might be that the pregnancy ends for a different reason, not because of the APS. This might be a chromosome problem, where something goes wrong very early in the baby’s development. In this case, the treatment for APS couldn’t help.

Either way, you may need time before you can begin to think about whether you would like to try again. And you might need extra support.

What about other health problems?

Now that you have been diagnosed with APS, your doctor should give you more information about other possible health risks related to blood-clotting problems. You will probably be advised to have regular check-ups at a specialist clinic in the future.

That might feel like a mixed blessing. All you were hoping for was an explanation for your miscarriages and a treatment to help prevent it happening again. Now you have to think not only about having a healthy pregnancy, but also about your general health.

It can be difficult to cope with this extra burden – indeed, you might feel it’s all very unfair. But it does mean that you will have the best possible care to help you and your baby in pregnancy and to help you stay healthy now and in the future.

Being told I had APS made me worry about my own health outside of pregnancy too. It was important to get specialist advice on things like long-haul flights.
Support and information

You might find it helps to talk to someone who can understand what you are going through.

All the organisations listed below offer support and information about pregnancy loss and/or APS. Your doctor may also find these useful sources of information.

The Miscarriage Association
Tel: 01924 200799
www.miscarriageassociation.org.uk
17 Wentworth Terrace, Wakefield WF1 3QW

The Miscarriage Association is a national charity which offers support and information on pregnancy loss (up to 24 weeks). You can talk to helpline staff or to a support volunteer who has experienced pregnancy loss and APS. You can also visit our online support forum.

Lifeblood: the thrombosis charity
Tel: 01558 650222
www.thrombosis-charity.org.uk
PO Box 58, Llanwrda SA19 0AD
Lifeblood works to raise awareness and promote research into thrombosis and offers impartial, factual information through its leaflets and on its website.

Hughes Syndrome Foundation
Tel: 020 7188 8217
www.hughes-syndrome.org
Lupus Unit, Gassiot House, St Thomas’ Hospital, London SE1 7EH
Offers support, information and education about Hughes syndrome as well as supporting research into the condition.

APEC – Action on Pre-eclampsia
Tel: 016 260 8088
www.apec.org.uk
2c The Halfcroft, Syston LE7 1LD
Offers support and information on pre-eclampsia through its helpline, leaflets and website.

SANDS – the Stillbirth and Neonatal Death Society
Tel: 020 7436 5881
www.uk-sands.org
28 Portland Place, London W1B 1LY
Offers support and information after stillbirth and neonatal death, through its helpline, leaflets and website.

“We all find different ways to cope.
I think talking to others on this forum helps you to not feel so alone.”

Need to talk to someone who understands?
Call our support line on 01924 200799. Monday to Friday, 9am-4pm
Or email info@miscarriageassociation.org.uk