menstrual cycle, convey concepts more dynamically than could ever be displayed on paper. Video consultations demonstrate more clearly our patients and people. The redemptive emphasis on the law, the client perspective and young people places contraception and sexual health within its philosophical, ethical and moral context. Links to referenced sites are well chosen and accessible. The interactive self-assessment is challenging and – dare I say – fun, and I learned from some errors but I will not confess where!

I think e-SRH e-Learning is a good preparation for the Practical Sessions of the FSRH Diploma, and with regular updating it will provide a valuable educational resource for us all in the future. Congratulations to all the team involved with this project.

Michael Topley, MBBS, DFFP
Associate Specialist, Contraception and Sexual Health, Choices Centre Towner Central, Stockport, UK.
E-mail: m.topley@bipenworld.com

References

Query about Faculty updated UKMÆK

I would be grateful if the Faculty of Sexual and Reproductive Healthcare could explain why in the updated UK Medical Eligibility Criteria for Contraceptive Use (UKMÆK 2009)1 the Category 4 for body mass index (BMI>40, has been removed? As a raised BMI is so closely associated with increased risk of venous thromboembolism, this does not seem logical. Without the Category 4 status, I am concerned that increasing numbers of patients with a BMI>35 and indeed a BMI>40, will start, or continue to take, the combined pill, without any robust guidance to support this as a dangerous practice.

I am, however, pleased to see the Category 3/4 for multiple risk factors for cardiovascular disease is now clearly stated. I would, however, prefer the definition for ‘older age’ to be stated. I would interpret this as being aged 35 years or over, but the additional comments at the end of the section imply the definition is aged 40 or above.

I really appreciate that UKMÆK is a guidance document and not a list of rules as such, but if these are too loosely presented then will they not serve their purpose in ensuring safe prescribing practice.

Deborah J Lee, MFFP, MRCGP
Associate Specialist in Reproductive Health, Lead Clinician CASH, Salisbury District of Sexual Health, Salisbury, Salisbury, UK.
E-mail: Deborah.J.Lee@salisbury.nhs.uk

Reference

Reply

In her letter, Dr Lee raises a pertinent question regarding the newly updated UK Medical Eligibility Criteria for Contraceptive Use (UKMÆK 2009)1 categories for body weight and combined hormonal contraception (CHC) use. The current Clinical Effectiveness Unit was not involved in updating UKMÆK but we believe the body weight categories were made less restrictive to make them more consistent with the categories for other cardiovascular risk factors and CHC.

The rationale for these changes is partly explained in an article by Trussell et al.2 Obesity is generally perceived to be an important risk factor in CHC users because of the high relative risk of venous thromboembolism (VTE). The article argues that, in terms of absolute or attributable risk, other cardiovascular risk factors are more strongly associated with VTE and mortality than obesity. For example, the absolute risk of VTE in CHC users aged 45–49 years (UKMÆK 2) is 175 per 100,000, which is greater than a VTE risk of 105 per 100,000 associated with CHC use and body mass index >35 (UKMÆK 3). The risks in terms of deaths in CHC users are even lower, with an absolute risk of 3.3 deaths per 100,000 in smokers aged <35 years (UKMÆK 2) and a risk of 0.8 deaths per 100,000 in women with BMI>35 (UKMÆK 3).

With regard to the UKMÆK 2009 section on multiple risk factors for cardiovascular disease, the text is unchanged from UKMÆK 2005. The additional comments do appear to imply that the UKMÆK definition of ‘older age’ is aged 40 years or above. Risk factors such as age are a continuum and there is not necessarily an exact cut-off. As Dr Lee acknowledges, UKMÆK is only a guidance document, and it would be entirely appropriate for clinicians to apply their own clinical judgement.

Louise Melvin, MRCOG, MFSRH
Director, FSRH Clinical Effectiveness Unit, and Consultant, Contraception and Sexual Health, Sandyn, Glasgow, G51 UK.
E-mail: louise.melvin@nhs.net

References

Implanon® failure in patients on antiretroviral medication: the importance of disclosure

We would like to draw other practitioners’ attention to a recently reported failure observed recently in our clinic, namely Implanon® failure in two women on antiretroviral (ARV) medication who failed to mention Implanon use to their HIV physicians. We wish to highlight the need for disclosure of HIV status prior to physicians offering contraceptive choices and Implanon use to the HIV physicians.

A 33-year-old woman, para 3, attended in May 2007, requesting termination of pregnancy. She had an Implanon since July 2004 and was amenorrhoeic until February 2007. She was commenced on Sustiva® (efavirenz 600 mg tab) and Truvada® (emtricitabine 200 mg and tenofovir disoproxil 245 mg) in January 2007 as her HIV viral load was rising. She was not asked and did not volunteer Implanon use. Concerns for the adverse effect of the ARVs on the fetus had prompted the termination request. After a normal dating ultrasound scan her decision to terminate became amiable. She was committed to the pregnancy after a 5-week scan and delivered a male baby weighing 3520 g at term.

A 35-year-old woman, para 1, conceived with an Implanon when commenced on efavirenz and lopinavir. She did not mention Implanon use until she visited the HIV physician, and the contraceptive clinic had no record of her HIV status. She did not appreciate that Implanon was a drug that might interact with ARVs. She described her Implanon and did not realise she was pregnant until 19 weeks. After counselling she opted for a termination.

The concentration of contraceptive hormones may change with concomitant drug use and vice versa. It is good practice to enquire about current and previous drug use (specifically lipid lowering enzymes) when offering hormonal contraception. With ARVs fulfilling some drugs might reduce hormonal contraceptive effectiveness. With the exception of the progestogen-only injectable or the levonorgestrel intrauterine system, the contraceptive efficacy of hormonal methods is reduced by liver enzyme inducers.

Some ARVs, such as protease inhibitors (atazanavir, ritonavir and non-nucleoside reverse transcriptase inhibitors) (efavirenz, nevirapine), are metabolised by the CYP3A4 liver enzyme system and can affect contraceptive efficacy.

Each Implanon contains 68 mg etonogestrel (ENG). The subdermal delivery method makes it 100% bioavailable. Serum ENG concentrations increase rapidly within 8 hours of insertion and peak after 4 days. The release rate is 60–70 μg/day in weeks 5–6 post-insertion, and decreases to 35–45 μg/day at the end of the first year, to 30–40 μg/day at the end of the second year, and then falls to 25–30 μg/day at the end of the third year.2 These low concentrations are sufficient to inhibit ovulation for 3 years.

The advice for using a hormone-only implant for women on long-term liver enzyme-inducing drugs is to continue using it together with additional contraceptive protection (such as condoms) and for 4 weeks after the drugs are stopped.

These cases highlight the unforeseen consequences of non-disclosure of HIV for both patients and physicians. One of the dilemmas facing physicians is whether to disclose the HIV diagnosis to general practitioners (GPs). Arguments have been advanced for specialists breaching confidentiality and notifying the GP against patients’ wishes in the interest of normal medical practice, the patients’ health personnel best interests, and the interests of society in general. Gillon4 examines each argument and concludes that none is sufficient to justify violating physician patient confidentiality in most cases.

Early contraceptive failure of Implanon in a woman on antiretroviral medication has been described.3 The patient in the case report had an ectopic pregnancy.

The majority of HIV-positive women are of reproductive age. Contraceptive options must take into account the risk of an unintended pregnancy, vertical transmission, and horizontal transmission for an HIV-positive partner. For HIV patients to achieve all these goals, a combined contraceptive (barrier method plus another method) is the ‘gold standard’. Some practitioners will argue that the ‘Double DUTCH’ advice should be given to all patients and not just HIV-positive women.

References