Migraine, the menopause and hormone replacement therapy: a clinical review

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Overview
In contrast to the relatively stable hormone profile during the middle reproductive years, the perimenopause is marked by erratic fluctuations of oestrogen and progesterone. By the late perimenopause, oestrogen levels can reach high magnitudes.1 Given the association between migraine and oestrogen ‘withdrawal’ it is not surprising that this stage of life is often accompanied by worsening migraine, which persists until oestrogen levels plateau post-menopause.

This paper reviews the effects that menopause and hormone replacement therapy (HRT) have on headache and migraine and considers the data on the risk of ischaemic stroke in HRT users with migraine. This evidence will be used to support practical guidance on the management of women with migraine wishing to use HRT.

Search strategy
A MEDLINE search from 1950 to April 2007 using the search terms ‘headache’, ‘migraine’, ‘menopause’, ‘hormone replacement therapy’, ‘perimenopause’, ‘progestogen’, ‘progesterone’, ‘(o)estrogen’, ‘(o)estradiol’ and ‘selective (o)estrogen receptor modulators’ identified 100 publications, which were scrutinised for relevancy to this review.

In addition, references from the author’s own files, a hand-search of the journals Cephalalgia and Headache, and peer-reviewed presentations at international congresses were considered.

Effect of the menopause on headache and migraine
Headache is a common but under-reported complaint in perimenopausal women. A study of 74 women attending a London menopause clinic reported that in the preceding 3 months, 57% had experienced headache and 29% had experienced migraine.2 Migraine was associated with significant disability, with 80% of women affected reporting attacks more often than once a month, 75% reporting severe attacks and 50% reporting attacks lasting longer than 1 day.

In 1000 women attending a Leicester menopause clinic, 85% reported recurrent headache of which 73% reported headache more often than once a month.3 Migraine was reported by 24% of women.

In a retrospective questionnaire of 29 postmenopausal women with tension-type headache, six women (21%) reported new onset of headache with menopause.4 Of those women who had had a physiological menopause, 27% reported improvement or complete remission of headache following menopause, 13% reported no change and 60% reported worsening headache. Of those women who had had a surgical menopause, 38% reported no change and 25% reported worsening headache.

Studies support the clinical impression that migraine without aura improves post-menopause, with time since menopause being a significant factor.5 A study of 1436 women at various stages of the menopause reported a prevalence of migraine of 10.5% in postmenopausal women following natural menopause compared with a prevalence of 16.7% in premenopausal and perimenopausal women [odds ratio (OR) 0.6, 95% CI 0.4–0.9, p = 0.03].6 This improvement is generally attributed to the absence of variations in sex hormone levels. In accordance with this, ovarian failure, with low levels of oestrogen and high levels of follicle-stimulating hormone, is associated with a lower prevalence of migraine than in menstruating women.7 However, psychological factors also seem to play a fundamental role.4

There are few data regarding the effect of menopause on migraine with aura, although the clinical impression is one of little change.

Type of menopause is important, with a lower prevalence of migraine following physiological menopause compared to surgical menopause (Table 1).4,7 Regarding the type of surgical procedures resulting in menopause, the prevalence of migraine has been reported as lowest in those with hysterectomy and bilateral oophorectomy, although not to a statistically significant level.2 Ovarian failure, with low levels of oestrogen and high levels of follicle-stimulating hormone, is associated with a lower prevalence of migraine than in menstruating women.7 Thus, it is not surprising that self-reported premenstrual syndrome was associated with a greater adverse effect of surgical menopause on migraine prevalence and a greater beneficial effect of natural ovarian failure.

The effects of medical menopause on migraine can be attenuated by the addition of HRT.8

Effect of HRT on headache
There are few data on the association between headache and current use of HRT. A retrospective questionnaire of 120 women attending a headache clinic in the USA suggested that HRT was associated with improved outcome, with 64.1% of respondents reporting improvement or complete remission of headache, 22.5% reporting no change and 13.3% reporting worsening headache.9 In contrast, a cross-sectional questionnaire of 6007 postmenopausal women taking part in large Norwegian health survey (Nord-Trøndelag Health Study) showed a significant association between headache and current use of HRT.10 This was irrespective of whether the route of delivery was local (OR 1.3, 95% CI 1.0–1.6) or systemic (OR 1.3, 95% CI 1.1–1.5).

Effect of HRT on migraine
Questionnaire studies suggest a significant association between migraine and current use of HRT (Table 2).10,11 Some studies suggest that a history of worsening migraine at menopause is a factor in predicting worsening migraine with HRT.3,9 However, it is not known whether HRT is associated with increased incidence of headache and migraine or whether HRT is initiated because of headache.


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Oestrogen
Regimen
Oestrogen replacement should be given continuously as oestrogen ‘withdrawal’ migraine can occur, in addition to return of menopause symptoms, with cyclical regimens.\textsuperscript{12}

Type of oestrogen
There are no data on the differential effects of human or equine oestrogens on migraine.

Route of delivery
Non-oral routes are less likely to have a detrimental effect on migraine than oral formulations of oestrogen replacement (Table 3).\textsuperscript{13,14} This is probably the result of the more stable serum hormone levels associated with non-oral routes.\textsuperscript{15} If oral preparations are favoured, tibolone may be the preferred option.\textsuperscript{16}

Dose
Oestrogen replacement therapy can have an adverse effect on migraine aura as evidenced by a report of 10 women seen in an ophthalmology clinic who were using transdermal oestrogen patches 50 µg daily, of whom six had a history of migraine (three migraine with aura, three migraine without aura) before using replacement therapy.\textsuperscript{17} All six women developed increased headache severity and accompanying visual scintillations. One patient with no previous history of migraine developed visual scintillations with no accompanying headache. Withdrawal of oestrogens and additional prophylactic antimigraine therapy led to marked improvement in all women, with complete cessation of migraine in four patients.

However, complete withdrawal of HRT may be unnecessary, as case reports on four women developing migraine aura following initiation of HRT showed that, in all cases, aura resolved with either a reduction in oestrogen dose or change in route of delivery of oestrogen.\textsuperscript{18}

Progestogen
Regimen
Continuous combined HRT appears to be better tolerated by migraineurs than cyclical combined HRT.\textsuperscript{19}

Type, route of delivery and dose of progestogen
There are no data regarding the effect on migraine of different types, routes of delivery or doses of progestogens used for HRT. However, progestogen intolerance resulting in bleeding problems, fluid retention, headache and negative mood adversely affects compliance.\textsuperscript{20}

<table>
<thead>
<tr>
<th>Table 1 The effect of menopause on migraine</th>
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<td>Study</td>
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<td>Neri et al. (1993)\textsuperscript{4}</td>
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<td>Granella et al. (1993)\textsuperscript{7}</td>
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<td>Mueller (2000)\textsuperscript{9}</td>
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<td>Martin et al. (2003)\textsuperscript{8}</td>
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NR, not reported.

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<th>Table 2 The effect of hormone replacement therapy on migraine: users versus never-users</th>
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<td>Study</td>
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<td>Misakian et al. (2003)\textsuperscript{11}</td>
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<td>Aegidius et al. (2007)\textsuperscript{10}</td>
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HRT, hormone replacement therapy; OR, odds ratio.
Migraine, the menopause and HRT

Table 3 The effect of different types of hormone replacement therapy on episodic tension-type headache and migraine

<table>
<thead>
<tr>
<th>Study design</th>
<th>Comparator drugs</th>
<th>Effect on ETTH</th>
<th>Effect on migraine</th>
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<tr>
<td>Nappi et al. (2001)</td>
<td>RCT of transdermal cyclical combined HRT vs oral cyclical combined HRT over 6 months in 30 women with migraine without aura and 20 women with ETTH</td>
<td>No change</td>
<td>No change</td>
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<td></td>
<td>50 µg transdermal oestradiol  patches daily plus 10 mg MPA on Days 15–28</td>
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<td>0.625 mg CEE daily plus 10 mg MPA on Days 15–28</td>
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<tr>
<td>Facchinetti et al. (2002)</td>
<td>RCT of oral continuous combined HRT vs oral cyclical combined HRT over 6 months in 38 women with migraine</td>
<td>Not applicable</td>
<td>Compared to baseline, treatment was associated in all three subgroups with increased in attack frequency (p &lt; 0.001), severity (p &lt; 0.001), days with headache (p &lt; 0.001) and analgesic consumption (p &lt; 0.001) although attack pattern were similar (p &gt; 0.05). Continuous combined regimen was better tolerated than cyclical and sequential regimens</td>
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<td>1 mg oestradiol hemihydrate plus 0.5 mg NET daily</td>
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<td></td>
<td>0.625 mg CEE daily plus 10 mg MPA on Days 15–28</td>
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<td>2 mg oestradiol valerate daily for 21/28 days plus 1 mg cyproterone acetate on Days 12–21</td>
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<td>Nappi et al. (2006)</td>
<td>RCT of low-dose oral continuous combined HRT vs oral tibolone over 6 months in 40 women with migraine without aura or ETTH</td>
<td>Reduced number of days with headache (p &lt; 0.005) Reduced severity (p &lt; 0.01) Reduced analgesic consumption (p &lt; 0.001)</td>
<td>Increased number of days with migraine headache (p &lt; 0.001) Increased number of analgesics (p &lt; 0.001)</td>
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<td>1 mg 17-beta-oestradiol plus 0.5 mg NET daily</td>
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<td></td>
<td>2.5 mg tibolone daily</td>
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CEE, conjugated equine oestrogens; ETTH, episodic tension-type headache; HRT, hormone replacement therapy; MPA, medroxyprogesterone acetate; NET, norethisterone acetate; RCT, randomised controlled trial.

Risk of cardiovascular disease in women with migraine

Migraine with aura is a risk factor for ischaemic stroke. Most of these studies assessed young women with ischaemic stroke and there are few data on older populations. More recent studies suggest that migraine with aura is also a risk factor for cardiovascular disease. Analysis of 5125 women aged over 45 years participating in the prospective cohort Women’s Health Study showed that only active migraine with aura at baseline, and not a past history, was a risk factor for ischaemic stroke, myocardial infarction, angina and death at 9 years follow-up. However, when migraine without aura was included, both conditions were associated with increased risk of cardiovascular disease and other major stroke risk factors.

Risk of cardiovascular disease in women using HRT

Concerns regarding an increased risk of cardiovascular disease associated with use of HRT are unjustified as they relate to an atypical population of postmenopausal women starting HRT. There is evidence from controlled trials to support earlier observations that HRT initiated during the perimenopause is not associated with increased risk of stroke and cardiovascular disease and is more likely to have beneficial effects.

On this basis there are theoretical benefits for women with migraine wishing to start HRT in the perimenopause. However, there are no data on risk of cardiovascular disease associated with new onset of migraine aura in women using HRT.

Risk of cardiovascular disease in women with migraine using HRT

There are few studies assessing the risk of ischaemic stroke or myocardial infarction in women with migraine using HRT. The Women’s Health Study did not find an association between these conditions and use of postmenopausal hormone therapy in women with migraine with or without aura.

Alternatives to HRT

Reviews and recommendations for effective alternatives to HRT in women with vasomotor symptoms support the use of fluoxetine, paroxetine, venlafaxine or gabapentin. Evidence of efficacy for migraine prophylaxis has been shown for fluoxetine and venlafaxine. Initial exacerbation of migraine in the first few weeks of treatment is an effect of some selective serotonin reuptake inhibitors (SSRIs) so it is important not to stop treatment.
too early. Gabapentin can also be used for migraine prophylaxis, although evidence of efficacy is far from robust. The most common adverse events reported are dizziness and sedation.

The efficacy and safety of complementary therapies have not been demonstrated, although phyto-oestrogens are widely used for vasomotor symptom control. One randomised controlled trial suggests some efficacy in menstrual migraine but, conversely, there is one case report of new-onset migraine following initiation of isoflavone supplements.

There are no data on the effect of selective oestrogen receptor modulators (SERMS) on postmenopausal women with migraine, although prescription-event monitoring has been used to assess safety in 13,987 patients for whom useful information was available. Flushing was the most common specific adverse event reported. Headache/migraine was the third most frequently reported adverse drug reaction associated with starting raloxifene and the fifth most frequently reported reason for stopping the drug. Although these data are limited, they do not support the use of SERMS as an alternative to HRT in women with migraine.

Practical recommendations for use of HRT (Box 1)

Headache in perimenopausal women is most likely to be a reflection of the background prevalence of migraine in the different age groups. Prospective data do not support an association between migraine with or without aura and ischaemic stroke in postmenopausal women, but do support an association between migraine with aura and increased risk of ischaemic stroke and myocardial infarction. Given the association between aura and cardiovascular risk factors, it is important to evaluate these risk factors and treat them appropriately.

There is no evidence that the treatment of menopausal symptoms in women with migraine should differ from standard recommendations, including use of HRT. There are theoretical and clinical benefits to non-oral oestrogen replacement for all women wishing to use HRT. For women with migraine, the evidence suggests that low-dose non-oral preparations of oestradiol should be recommended as first choice. Continuous progestogens are better tolerated than cyclical progestogens. Changing the type of progestogen can help, as side effects are fewer with progestrone derivatives such as medroxyprogesterone acetate and dydrogesterone rather than with testosterone derivatives such as norethisterone. Drospirenone is a more recent well-tolerated option although it is currently only available as an oral preparation combined with oestrone. Changing route from oral to transdermal progestogen may also be effective, as may reducing the course of cyclical progestogens to only 7–10 days per month, although the latter is potentially associated with increased risk of endometrial hyperplasia. Natural progesterone is available as suppositories, micronised tablets and vaginal gel, although the availability of these different formulations varies worldwide. Sedation is a common adverse effect of natural progesterone. A well-tolerated option is the levonorgestrel-releasing intrauterine system used with oestradiol supplementation.

Occasionally progestogenic adverse effects are sufficiently bothersome for a woman to choose to discontinue progestogens. In these cases specialist referral is appropriate for regular endometrial assessment.

Although migraine attacks may increase when HRT is initiated, there is usually improvement with continued use. Any new-onset headache should be carefully evaluated for secondary causes, as although migraine may occasionally develop for the first time during the perimenopause, it is unusual for migraine to begin post-menopause.

There are no data regarding risk associated with developing a first attack of migraine with aura when using HRT. There is the concern that transient ischaemic attacks (TIAs) may be misdiagnosed as aura since it is not always easy to distinguish between the two conditions. On a practical basis, once TIAs have been excluded, the dose and route of delivery of oestrogen replacement should be assessed to provide the lowest effective dose necessary to control menopausal symptoms.

For symptomatic and prophylactic treatment of migraine standard strategies apply. Given the association between migraine aura and increased risk factors for cardiovascular disease, these should be assessed and managed accordingly. Unless the woman’s cardiovascular risk profile contraindicates triptans, there is no evidence that triptan use is associated with increased risk of stroke or cardiovascular disease.

Conclusions

Headache and migraine are common symptoms of the menopause but are often under-reported. Perimenopausal women should routinely be asked about headache and migraine so that they can be offered appropriate advice. This should include optimal symptomatic treatment for attacks and strategies for prevention. In women with additional menopausal symptoms, management may include use of continuous HRT via a non-oral route, using the lowest effective dose necessary to control symptoms.

Box 1: Optimising hormone replacement therapy (HRT) for women with migraine

- Migraine aura is not a contraindication to use of HRT.
- If aura appears for the first time after starting HRT, reduce the dose of oestrogen and consider changing route of delivery (oral to patch, patch to gel).
- Use the lowest effective dose of non-oral oestrogens that will control vasomotor symptoms.
- Where progestogen is required, continuous delivery is best, with preparations such as:
  - levonorgestrel intrauterine system
  - progestosterone-derived progestogens (e.g. dydrogesterone) rather than testosterone derivatives (e.g. norethisterone)
  - drospirenone
  - micronised progesterone, or progesterone suppositories or vaginal gel, if available.
- If progestogens are not tolerated, consider discontinuation, but with regular endometrial assessment.

References

Migraine, the menopause and HRT


