SHORT REPORT

Dose and duration of hormone use: understanding the effects of combined menopausal hormones on breast cancer better, 1976–2004

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The effects of exogenous hormones on diseases that are hormonally related, particularly breast cancer, cause concerns. Breast cancer is common and the relation with hormones complex. Millions of women have taken menopausal oestrogen and progesterone (hormone therapy, HT) for prolonged periods to alleviate menopausal symptoms, to prevent osteoporosis, and to prevent coronary heart disease too.

The breast cancer risk of HT has been investigated since the 1970s in some 50 studies. It has been known that it was associated with an increase in breast cancer, but there has been confusion because of the possible differential effect of different hormones as well as of different durations, methods of application, and the effect of past use. It was none the less argued that the risk was small. It is surely a public health responsibility to clarify these risks as far as possible. This is particularly so when the marketing imperatives are so strong.

Use of combined therapy became common in the 1980s, after years of use of oestrogen alone were found to increase endometrial cancer risk, which the addition of progesterone eliminated. Thus epidemiological data on combined therapy were less mature and often confused with data on unopposed therapy, which had been the subject of most studies. The addition of progesterone could, in theory, have had a differential effect on breast cancer risk in any direction.

The million women study, published in 2003, showed that five years or more of use of combined HT doubles the risk among current users, which is more extreme than oestrogen alone. Such a risk has important public health implications, when 25% of menopausal women were estimated to use HT by 2000. How much could we have known, by when, of the effect of long term combined HT use, with better pharmacovigilance? To investigate this question we have used quick and efficient statistical methods that require simple analyses of published data, and thus were available to all.

METHODS

We undertook a systematic review of the published research literature of all studies that examined the relative risk of breast cancer with five or more years of recent combined HT use. Some studies did not distinguish between unopposed or combined use and we included them only if the proportion of combined use was stated as being 30% or more. We pooled the risk estimates; using conventional fixed effect meta-analysis, calculated cumulative pooled estimates after each eligible study was published (final three columns in table 1). Where studies reported risk for several categories of use over five years we similarly pooled these estimates to provide one weighted estimate for each study. Relative statistical weights were estimated from the stated confidence limits of odds ratio (OR) estimates. Also where risks were only reported separately for subgroups, we pooled the estimated effects for the category closest to current use of five or more years of combined HT.

RESULTS

It may be seen in table 1 that the million women study dominates any final estimates statistically as it was so large. It however estimates higher risks (significantly) than the synthesis of all other previous studies. Some small individual studies estimate risk higher than 2.0, but this is to be expected by chance. However by 1989 a pooled relative risk of around two was significant and hence should have been

| Table 1 | Cumulative synthesis of all studies examining the risk of breast cancer with more than five years use of combined HT, by publication year |
|---|---|---|---|---|---|---|
| | Study | Year | % Comb (adj) | OR | lower CI | upper CI | weight |
| Hunt | 1987 | 43 | 3.60 | 0.90 | 15.00 | 2 |
| Ewertz | 1988 | 35 | 1.75 | 1.30 | 2.36 | 43 |
| Bergkrist | 1989 | 100 | 4.40 | 0.90 | 22.40 | 2 |
| Harris | 1992 | 39 | 1.20 | 0.65 | 2.23 | 10 |
| Newcomb | 1995 | 100 | 1.12 | 0.72 | 1.76 | 20 |
| Stanford | 1995 | 100 | 0.56 | 0.32 | 0.97 | 12 |
| Magnusson | 1999 | 100 | 2.46 | 1.97 | 3.08 | 78 |
| Shailer | 2000 | 100 | 1.74 | 1.22 | 2.48 | 30 |
| Ross | 2000 | 100 | 1.63 | 1.43 | 1.85 | 224 |
| Celbowski WHI | 2003 | 100 | 1.97 | 1.36 | 2.85 | 28 |
| Li | 2003 | 100 | 2.20 | 1.64 | 2.95 | 45 |
| Overview | 1997 | 100 | 1.53 | 0.88 | 2.18 | |
| Beral MWS | 2003 | 100 | 2.21 | 2.06 | 2.36 | 778 |
| Cumulative meta-analysis | | | | | | |
| Cumulative pooled ORs | | | | | | |
difficult to ignore. Publication bias may be in part responsible as the pooled estimate decreases during the 1990s, attributable to two studies with low risk estimates. Possibly this suggests the intrinsic unreliability of observational epidemiological study of a heterogeneous product, with changing use patterns. There is some heterogeneity between all these studies in both exposure and estimated effect, giving rise to opportunities for varied interpretations, but the net effect is unambiguous.

The large overview from the individual records from 51 studies published in 1997 shows a relative risk that is entirely consistent with this more simple synthesis of published results. There was no deemed to be little use of combined preparations in the collaborations studying, resulting in no significant effect of five or more years of combined HT use in 1997. The simpler analyses reported here show a significant effect well before that.

CONCLUSIONS AND RECOMMENDATIONS

The particular subgroup of exposure investigated was suggested partly from the results of recent studies. However, it does not seem excessive to suggest that any pharmaceutical product that has the potential for widespread use should be routinely investigated for possible serious long term side effects with respect both to essential composition (both constituent and dose) and duration of use.29 In this case the opportunity for systematically testing essentially two products (unopposed and combined HT) and two durations (less than five years and more than five years) for long term side effects from both randomised and observational studies was missed both by the public health community and regulators. But the ultimate responsibility rests with the manufacturers and their marketing departments—none the less independent assessment is essential. In this case the potential danger of long term use of combined HT on breast cancer risk could have been identified much earlier—but the pressure to accept any reliable notes of caution were powerless resisted unless and until the evidence became entirely convincing, because of the real and perceived benefits of HT. In particular its putative cardioprotective effects were frequently cited—until shown to be illusory.20

In the UK around 15 000 diagnoses of breast cancer are made each year among women aged 50–64. Acting on these results by using combined therapy sparingly strictly for symptoms, because of the increased breast cancer risk, might thus have saved around 2000 of these women from that disease per year. Clearly there is an issue with the downward trend in the early 1990s that would have been difficult to interpret without the benefit of hindsight. But since 1989 the pooled relative risk estimate was never non-significant, mostly highly significant and close to two.

The long term safety of medicinal products designed to alleviate common symptoms in the short term and that therefore have an enormous potential market require rigorous and routine assessment. Epidemiological studies should always report findings separately for drug type and categorised into meaningful duration groups. The paucity of epidemiological data in some groups early in the experience of a product is not a special problem when subjected to proper meta-analysis across studies as they accumulate. Indeed it is an essential part of such assessment. But it is necessary to be rigorous about sensible subgroups of exposure and outcome, a priori. This should be routine.

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