Change of seizure frequency in pregnant epileptic women

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SUMMARY The effect of pregnancy on seizure frequency was monitored prospectively in 136 pregnancies of 122 epileptic women. Pregnancy did not influence the seizure frequency in 68 pregnancies (50%). In 50 pregnancies (37%) the number of seizures increased during pregnancy or puerperium. The seizure frequency decreased in 18 pregnancies (13%). In 34 out of 50 pregnancies (68%) the increase was associated with non-compliance with the drug regimen or sleep deprivation. In seven out of 18 pregnancies (39%) improvement was related to correction of non-compliance or sleep deprivation during the pregestational nine months. Insufficiently low plasma concentrations of antiepileptic drugs were found in 47% of the women with uncontrolled epilepsy during pregnancy. The course of epilepsy during pregnancy is primarily influenced by non-compliance, sleep deprivation during pregnancy, and inadequate therapy before and during pregnancy. With good medical attention pregnancy itself seems to have only a minimal influence on the course of epilepsy.

Improvements in the drug therapy of epilepsy have led to a growing number of well controlled epileptic women in childbearing age. It is therefore not surprising that the influence of pregnancy on the course of epilepsy has received renewed interest. According to a recent review, the seizure frequency improved in 23% and deteriorated in another 24% of 2165 pregnancies.1 A number of variables may influence the seizure frequency. These include hormonal and metabolic factors, respiratory changes, psychological problems, non-compliance with the prescribed drugs regimen, changes in pharmacokinetics of antiepileptic drugs, and a modification of seizure propensity during pregnancy.1

Preliminary studies suggested that non-compliance, seizure provocation, and a decrease in the plasma concentration of antiepileptic drugs may be primarily involved in the regulation of seizure frequency during pregnancy.2,3 Changes in seizure propensity can be studied directly in pregnant women not receiving antiepileptic drugs. The purpose of this prospective two-centre study of 136 pregnancies was to investigate changes in seizure frequency during pregnancy and to analyse clinical and pharmacological factors associated with a seizure relapse in the individual patient.

Patients and Methods

A total of 136 pregnancies of 122 epileptic women were followed prospectively from 1977 to 1981 in West-Berlin, Germany and Milan, Italy. Pregnant women were excluded from the study if they had a total of less than three verified epileptic seizures, the data on the seizure frequency were incomplete or the pregnancy was ended by abortion. The patients were referred to the Epilepsy Clinic in Milan and Berlin from departments of gynaecology and neurology and gynaecologists and neurologists for pregnancy planning and during early pregnancy.2,3 The project was made known to them by regular distribution of information in letters and reprints. The patients were evaluated neurologically and usually seen at monthly intervals, when the history of seizures was recorded. At each visit the patients were examined for clinical drug toxicity. Special emphasis was given to the history of the clinical factors possibly associated with an increase in seizure frequency or a relapse of seizures. The patients were asked specifically

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whether they had taken the daily dose regularly in the days before each individual seizure. Non-compliance was assumed only when the patient admitted to it. Furthermore, information was sought on seizure provocation by sleep deprivation, and consumption of ethanol, or both. Sleep deprivation was defined as a delay of more than 2 hours from the usual working day onset of sleep for the individual patient. Sleep deprivation was assumed to be a predisposing factor only in patients with a previous history of seizure provocation through sleep deprivation. At each visit the plasma concentration of phenytoin, phenobarbitone, carbamazepine, ethosuximide and valproic acid was measured in duplicate by enzyme immunoassay as described previously. For quality control the laboratories joined the European Quality Control Scheme organised by Prof A. Richens.

One drug treatment was administered in 86 pregnancies (63%) with carbamazepine (n = 12), primidone (n = 22), phenobarbitone (n = 20), phenytoin (n = 23), valproic acid (n = 4), ethosuximide (n = 3) or clobazam (n = 2). Two antiepileptic drugs were prescribed in 25 pregnancies and three drugs were given during two pregnancies. During 23 pregnancies the epileptic women did not take any antiepileptic drug. The clinical data of this series are given in table 1. For evaluation of the course of epilepsy, the number and type of seizure was compared in each patient during the nine months before pregnancy and during each trimester of pregnancy. We also analysed changes in seizure frequency during the puerperium that is the three months following the delivery. Fifty-three patients were followed up to the 12th month following the pregnancy. Increase or decrease of seizure frequency was defined by giving the actual seizure frequency. Percentages alone were considered inadequate as a means of defining change.

Results

Pregnancy did not influence the seizure frequency in 68 pregnancies (50%). In 38 of the 68 pregnancies the epileptic women had had no seizures in the pregestational nine months and during pregnancy. In 50 pregnancies (37%) the number of seizures increased during pregnancy or puerperium. Finally, the number of seizures decreased during pregnancy in 18 pregnancies (13%).

Increased seizure frequency

The number of seizures increased in 50 pregnancies (table 2). Non-compliance or sleep deprivation were noted in a temporal relationship to a seizure relapse in 34 out of 50 pregnancies (68%). Except for a higher incidence of generalised epilepsy as compared to focal epilepsy (59% vs. 13%) (p = 0.02), the non-compliers did not differ in clinical characteristics from the other patients. Non-compliance or sleep deprivation were implicated in 27 out of 34 pregnancies (79%) with an increase of generalised tonic-clonic seizures and in five out of 12 pregnancies (42%) with increased simple or complex-partial seizures. In 29 pregnancies (58%) the increase was first seen during the first trimester (fig 1). The increase in seizure frequency during puerperium was related to non-compliance or sleep deprivation in five of six pregnancies.

Table 1 Clinical features of the patients

<table>
<thead>
<tr>
<th>Clinical data of the patients</th>
<th></th>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No of patients</td>
<td>122</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of pregnancies</td>
<td>136</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at delivery</td>
<td>25 ± 4 years</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Generalised epilepsy</td>
<td>60</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Focal epilepsy</td>
<td>62</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration of epilepsy</td>
<td>13.7 ± 6 years</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of patients with a total of less than seven generalised tonic-clonic seizures</td>
<td>42* (39%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of pregnancies with one or more seizure types</td>
<td>51</td>
<td>78</td>
<td>7</td>
<td></td>
</tr>
</tbody>
</table>

*No information in six patients, no generalised tonic-clonic seizures in nine patients, resulting in 107 patients.

Table 2 Type of seizures, non-compliance and sleep deprivation in pregnancies with increased seizure frequency

<table>
<thead>
<tr>
<th>Type of seizures</th>
<th>Total</th>
<th>NC-SD</th>
<th>No seizure before pregnancy</th>
<th>Seizures before pregnancy</th>
<th>Percent increase in seizures* (mean and range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GTC alone</td>
<td>34</td>
<td>27</td>
<td>15</td>
<td>14</td>
<td>+233% (100-400) (8)a</td>
</tr>
<tr>
<td>GTC and CPS</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>+100 and 200% (1)b</td>
</tr>
<tr>
<td>GTC and SPs</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>CPS or SPs</td>
<td>12</td>
<td>5</td>
<td>7</td>
<td>4</td>
<td>+270% (255-300) (3)c</td>
</tr>
<tr>
<td>Absence</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>nd (2)</td>
</tr>
</tbody>
</table>

GTC = generalised tonic-clonic seizures; CPS = complex-partial seizures; SPs = simple partial seizures; NC = non-compliance in 15 women; SD = sleep deprivation in 14 women, and both, NC and SD, in 1 woman; * = No of patients with this type of seizure in the nine months before pregnancy; nd = not determined as quantification of absence seizures was difficult.

Actual seizure frequency of the individual patient in the nine months before and during pregnancy:

a = GTC: 1 vs. 2, 1 vs. 3, 1 vs. 4, 9 vs. 44, 5 vs. 12, 17 vs. 40, 4 vs. 20, 9 vs. 20
b = GTC: 2 vs. 4 and 1 vs. 3 cps,
c = cps: 2 vs. 8, 11 vs. 39, 9 vs. 32
Change of seizure frequency in pregnant epileptic women

The onset of increased seizure frequency in the three trimesters of pregnancy (1, 2, 3), and during puerperium (p). The figure shows that in most pregnancies the increase begins in the first trimester, while it is lowest in the third trimester. Non-compliance and sleep deprivation account for 5 out of 6 pregnancies (83%) with increased seizure frequency during puerperium.

The rate of increase varied widely. Thirteen out of 23 patients who were completely free of seizures in the nine months before pregnancy developed a single generalised tonic-clonic seizure and two patients had more than one generalised tonic-clonic seizure. Seven patients had three to six simple or one to three complex-partial seizures, and one patient had a single generalised tonic-clonic seizure and simple focal seizures. Among the 27 patients who were uncontrolled in the nine months before pregnancy, 13 patients developed either one to three generalised tonic-clonic seizures or two complex-partial seizures which they did not have in the nine months before pregnancy. In the other 14 patients, the average increase in percent was 223% for generalised tonic-clonic seizures and 270% for simple or complex-partial seizures (table 2).

Antiepileptic drug monitoring could be evaluated in 11 out of 16 pregnancies in which non-compliance or sleep deprivation was not involved in seizure precipitation. In nine pregnancies of women with previously uncontrolled epilepsy the plasma concentration was not increased or fluctuated widely during pregnancy. In one pregnancy a decrease in plasma concentration was not associated with an increased seizure frequency, while only in one out of 11 pregnancies the number of seizures increased despite a higher plasma concentration.

Decreased seizure frequency

The pregnancy seemed to have a beneficial effect on seizure frequency in 18 pregnancies. The number of seizures decreased by 50% (23%-70%) in seven patients, and 11 patients became completely controlled during pregnancy. In seven women non-compliance or sleep deprivation was associated with seizures in the gestational nine months, while the women remained free of seizures during pregnancy. In four out of the remaining 11 pregnancies the plasma concentration was increased or a new drug

<table>
<thead>
<tr>
<th>Clinical features</th>
<th>Change in seizure frequency during pregnancy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No change</td>
</tr>
<tr>
<td></td>
<td>n</td>
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<tr>
<td>Age at the delivery ≥ 28 years</td>
<td>7</td>
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<tr>
<td>Primary generalised epilepsy</td>
<td>42*</td>
</tr>
<tr>
<td>Focal epilepsy</td>
<td>26†</td>
</tr>
<tr>
<td>One type of seizures</td>
<td>28‡</td>
</tr>
<tr>
<td>Two types of seizures</td>
<td>36</td>
</tr>
<tr>
<td>Three types of seizures</td>
<td>4</td>
</tr>
<tr>
<td>Duration of epilepsy ≥ 10 years</td>
<td>44</td>
</tr>
<tr>
<td>No generalised tonic-clonic seizure in the year before pregnancy</td>
<td>49§</td>
</tr>
<tr>
<td>Total or less than seven generalised tonic-clonic seizures</td>
<td>28</td>
</tr>
<tr>
<td>Family history of epilepsy</td>
<td>6</td>
</tr>
</tbody>
</table>

*p = 0.07
†p = 0.06
‡p = 0.08
§p = 0.05

[11 patients were seizure free during pregnancy; in 7 patients, the actual seizure frequency in the nine months before and during pregnancy were: CPS: 90 vs. 45, 36 vs. 19, 5 vs. 2; GTC: 4 vs. 1, 6 vs. 2, 30 vs. 23; GTC: 70 vs. 21 and 45 vs. 20 cps.]
was introduced during pregnancy. In five pregnan-
cies the plasma concentration remained unchanged
during pregnancy. During two pregnancies no drug
treatment was given.

We found no clinical features which help to pre-
dict an increase or a decrease of seizure frequency in
the individual patient (table 3). Primary generalised
epilepsy, patients with only one type of seizure and
no generalised tonic-clonic seizure in the year
before pregnancy were fewer in the group with an
increase in seizure frequency. The difference did not
reach statistical significance of p < 0.05, however
(table 3).

In 53 pregnancies the number of seizures was
compared from nine months before to 12 months
following the pregnancy. In 34 pregnancies (64%) a
similar number of seizures or absence of seizures
was recorded in the pregestational nine months and
during months 4 to 12 following the delivery. In 17
pregnancies the seizures returned to the pregesta-
tional frequency after an increase or a decrease dur-
ing pregnancy. In 19 pregnancies the number of
seizures was lower (n = 17) during the pregesta-
tional period, and only rarely higher (n = 2).

The course of epilepsy was studied during 23
pregnancies without antiepileptic drug therapy.
There was no change in seizure frequency in 13 out
of 23 pregnancies (57%). In eight pregnancies
(35%) the number of seizures increased. Sleep
deprivation was responsible in six of eight pregnan-
cies. In two pregnancies (9%) the number of seiz-
ures improved. When compared to the changes in
seizure frequency in women receiving drug treat-
ment there was no significant difference.

The first antiepileptic drug plasma concentra-
tion obtained during pregnancy was either below 10
µg/ml phenytoin, or 25 µg/ml phenobarbitone 4
µg/ml carbamazepine or 50 µg/ml ethosuximide and
valproic acid in 63% of all 98 pregnancies despite
uncontrolled seizure before (n = 75) or during
(n = 23) pregnancy. The highest plasma concentra-
tions of antiepileptic drugs during pregnancy were
still below this range in 47% of the patients with
uncontrolled epilepsy.

Discussion

Pregnancy had either no or a beneficial influence on
the seizure frequency in nearly two-thirds of all
pregnancies. In 50 pregnancies (37%) the number
of seizures increased during pregnancy. This overall
result is similar to that of prospective studies in Fin-
land and Canada, and retrospective reports.

A discussion of the factors responsible for a
change in the course of epilepsy must consider the
random fluctuations of seizure frequency observed
during a 18 months period in any patient regard-
less of pregnancy, and try to establish factors specific for
the pregnant epileptic women which could possibly
influence the course of epilepsy. At the onset of
pregnancy the seizures were uncontrolled in 75
pregnancies (55%) suggesting less than optimal
drug treatment prior to the pregnancy if one accepts
that complete seizure control can be achieved in
70–80% of the patients with adequate drug
therapy. This is supported by the finding that 63%
of these women were treated with inadequately low
plasma concentrations of antiepileptic drugs. The
highest plasma concentrations of antiepileptic drugs
during pregnancy were still below the accepted
therapeutic range in 47% of the pregnant women
with uncontrolled epilepsy.

One mechanism for a reduction of the seizure fre-
quency during pregnancy was the improvement of
inadequate therapy prior to the pregnancy or the
explanation of non-compliance and sleep deprivation
in 11 of 18 pregnancies (61%). In seven pregnan-
cies the improvement of seizure frequency cannot be
explained and may be due to random fluctuation of
seizure frequency or other factors (for example
hormonal changes). The finding that an improved
drug treatment, and not the pregnancy itself, may be
responsible for a reduction of seizures suggests that
the reverse mechanism may be partly responsible for
a deteriorated course of epilepsy during pregnancy.
In fact, in 34 out of 50 pregnancies (68%) admitted
non-compliance with the prescribed drug regimen or
sleep deprivation were associated with an increase in
seizure frequency. This preventable mechanism of
seizure provocation is therefore a previously under-
estimated major factor which determines the course
of epilepsy in pregnancy. Non-compliance or sleep
deprivation may be suspected in a woman with
generalised epilepsy, a single relapse of mostly
generalised tonic-clonic seizures, and a history of
non-compliance or seizure provocation through
sleep deprivation. Non-compliance with the drug
regimen is most frequent at the onset of pregnancy,
usually when the woman finds out that she is
pregnant. Fear of malformations or other adverse
effects of antiepileptic drugs are given by a number of
women as the motivation to stop the treatment.
Unfortunately, some physicians supported the
women in their intent to reduce or stop the drug
treatment, possibly because of the impression that
the teratogenic risk increases with a higher dose of
the antiepileptic drug. Currently, there is no evi-
dence to support this impression.

Of further concern is the finding that the relapse
of seizures during the puerperium is associated with
either an insufficient drug therapy, that is the patient
does not receive her daily dose because the obstetri-
cian is unaware of her drug treatment or someone has simply forgotten to give her the drug in the hospital. Additional sleep deprivation which may occur throughout the pregnancy, is characteristically associated with a relapse of seizure during breast feeding in the puerperium. Care should be taken that the mother has her regular hours of sleep, possibly through the help of the husband or a relative who takes care of the child at night. In 16 out of the 50 pregnancies with increased seizure frequency non-compliance or sleep deprivation were not involved according to the patient’s history. In nine of these pregnancies the drug treatment was not sufficient to increase the steady-state plasma concentration above those during pregestational period where the patients were uncontrolled. In one pregnancy the seizure frequency increased despite higher plasma concentration during pregnancy. Consequently, the increase in seizure frequency remains unexplained in six pregnancies and may entirely reflect the spontaneous course of chronic epilepsies during a period of 18 months regardless of pregnancy. Furthermore, sleep deprivation was responsible for the increased seizure frequency in six of eight pregnancies of women not taking antiepileptic drugs indicating that it is a major factor for the deterioration of untreated epilepsies. In this series clinical data other than those related to non-compliance and sleep deprivation were not helpful to predict the individual’s course of epilepsy during pregnancy. We could not confirm the value of the pre-pregnancy frequency of generalised tonic-clonic seizures, the duration of epilepsy, and the course of epilepsy in previous pregnancies as suggested by Remillard et al. This report is in agreement with the large Finnish series of Bardy who likewise found no clinical factors which predict the increase in the number of seizures. The clinical implication of our data for the management of the pregnant epileptic woman suggest that a renewed effect should be made to completely control the epilepsy with the help of adequate drug therapy when pregnancy is planned. Specific risks include the discontinuation or reduction of drug therapy especially during early pregnancy, deficient drug delivery in the obstetric department, and seizure provocation through sleep deprivation during pregnancy and breast feeding.

In conclusion then, pregnancy itself does seem to have only a minimal effect on the course of epilepsy, if at all. In the few patients with an unexplained increase of seizure frequency the influence of other factors (e.g. hormonal, random fluctuations of seizure frequency) remains to be evaluated.

References


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