Molecular genetic evidence for unifocal origin of advanced epithelial ovarian cancer and for minor clonal divergence

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Summary Detection of loss of heterozygosity (LOH) and DNA flow cytometry (FCM) were used to trace the origin of bilateral ovarian cancer from 16 patients. From each tumour the DNA index (DI) and LOH patterns for chromosomes 1, 3, 6, 11, 17, 18, 22 and X were determined with 36 microsatellite markers. Formalin-fixed, paraffin-embedded as well as frozen specimens were used. Flow cytometric cell sorting was used to enrich tumour cells for polymerase chain reaction (PCR)-driven LOH analysis. Analysis of the LOH data showed that in 12 of the 16 cases concordance was observed for all informative markers, namely retention of heterozygosity (ROH) or loss of identical alleles in both tumour samples. In four cases discordant LOH patterns were observed. In two cases the discordant LOH was found for one of the chromosomes tested while other LOH patterns clearly indicated a unifocal origin. This suggests limited clonal divergence. In the other two cases, most likely independent origins were observed. The number of chromosomes showing LOH ranged from 0 to 6. Comparison of DNA FCM and the LOH data showed that the latter technique has a higher sensitivity for the detection of a unifocal origin. In 14/16 cases evidence was found for a unifocal origin, while in two cases clonal divergence was found at LOH level and in two other cases clonal divergence at DNA ploidy level. In 12 cases the complete observed alleleotype had developed before the formation of metastases, including the two cases showing a large DNA ploidy difference.

Keywords: cell sorting; clonality; DNA flow cytometry; paraffin-embedded tissue

Epithelial cancer of the ovary has the poorest prognosis of all gynaecological cancers with a 5 year survival of 39% (Boring et al., 1991). More than 50% of these malignant ovarian tumours are bilateral (Novak and Woodruff, 1974) and frequently tumour locations are found in other sides of the gynaecological tract as well. It is often uncertain whether they represent metastatic disease or multiple primary tumours, since histopathological characteristics are often not sufficient to solve this dilemma.

Pejovic et al. (1991) studied this problem by comparative karyotype analysis and found identical chromosome changes in 15 of the 18 cases for which multiple tumour deposits were studied, although large karyotypic differences between the cases were observed. In three cases no karyotypic abnormalities were observed. Smit et al. (1990) used DNA FCM to compare DNA indices (DIs) of multiple tumours. In 60% of the cases the similarity in DI provided evidence for metastatic disease. In addition molecular genetic studies provided evidence against a multifocal origin. Mok et al. (1992) compared multiple tumours of the same patient for their p53 mutation pattern. In the nine cases investigated the mutations were identical in both tumour sites. Jacobs et al. (1992) compared tumours on the basis of LOH at different chromosomes, p53 expression, p53 mutations and X chromosome inactivation pattern. Statistical analysis showed that the probability of an independent occurrence of the tumours was less than 1.5% in 15 of 17 cases.

In the present study we have investigated the clonality of bilateral ovarian carcinomas by DNA FCM and molecular genetic analysis with particular attention to possible clonal divergence after metastasis. To this end we used 36 polymorphic microsatellite markers mapping to eight different chromosomes for LOH analysis. Furthermore, we compared the discriminating power of DNA FCM as a rapid technique with the more laborious approach of LOH analysis. To eliminate the interfering effect of contaminating normal DNA, we applied flow cytometric cell sorting on the basis of DNA content to enrich tumour cell populations for PCR-driven LOH analysis.

Materials and methods

Specimens

Sixteen cases of epithelial ovarian cancer with bilateral tumour locations were used (Table I). Nine were formalin-fixed, paraffin-embedded archival specimens and seven were snap-frozen fresh specimens stored at −70°C. Tumour cell percentages were determined by visual examination of haematoxylin- and eosin-stained sections. When specimens contained less than 80% tumour cells, they were enriched by flow cytometric cell sorting (Table I) based on DNA ploidy as described below. Constitutional DNA was extracted from peripheral blood leukocytes. If blood was unavailable, archival paraffin blocks with normal (non-tumour) tissue from the same patient or flow cytometrically sorted, DNA diploid, stromal cells from the aneuploid tumour served as the source for constitutional DNA (Table I).

Flow cytometry and cell sorting

Suspensions of paraffin-embedded nuclei were obtained according to Hedley et al. (1983) with minor modifications (Schueler et al., 1993) and stained with propidium iodide after RNAse treatment. Fresh frozen specimens were processed according to the detergent-trypsin method of Vindelov et al. (1983a). Trout erythrocytes were used as reference (Vindelov et al., 1983b). Samples were analysed on a FACScan flow cytometer (Becton Dickinson, Mountain View, CA, USA). Flow cytometric sorting was performed on the basis of DNA content on a FACStar flow cytometer (Becton Dickinson). DIs were calculated according to accepted criteria (Hiddemann et al., 1984).

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DNA isolation

DNA was extracted from peripheral blood lymphocytes according to Miller et al. (1988). DNA from fresh tumour specimens was isolated as described by Devilee et al. (1989). DNA from paraffin-embedded tissue was isolated by overnight incubation of six 10 µm sections in 100 µl PK buffer [10 mmol l⁻¹ Tris-HCl pH 8.3, 0.5% Tween 20, 1 mmol l⁻¹ EDTA 1% proteinase K (w/v)] (Limpens et al., 1993), at 56°C followed by a 10 min incubation at 100°C to inactivate proteinase K. To obtain a proper separation of the liquid and solid phases, the tubes were centrifuged directly while samples were hot. The liquid phase was carefully pipetted through the solid paraffin residue laying on top of the sample. DNA was isolated from the sorted nucleus fractions as described before (Abeln et al., 1994). Briefly, nuclear concentrations were adjusted to 50 µg per µl with PK buffer and incubated overnight at 56°C followed by a 10 min proteinase K inactivation at 100°C.

Detection of LOH

The microsatellite markers that were selected on the basis of heterozygosity and chromosomal location are presented in Table II. At least one marker was used for each chromosome, depending on interpretability and informativeness of the alleles. In non-informative cases, adjacent markers were used. Analysis of LOH was performed by PCR on microsatellite markers as described by Weber and May (1989). Briefly, 2 µl of DNA solution was loaded on a microtitre plate. Reaction mix was added to a total reaction volume of 15 µl containing 3 pmol of the forward and 3 pmol of the reverse primer, 1% bovine serum albumin (BSA), 0.01% gelatin, 0.1% Triton X-100, 10 mmol l⁻¹ Tris-HCl pH 9.0, 50 mmol l⁻¹ potassium chloride, 1.5 mmol l⁻¹ magnesium chloride, 200 mmol l⁻¹ dATP, 200 mmol l⁻¹ dGTP, 200 mmol l⁻¹ dTTP, 0.25 mmol l⁻¹ dCTP and 0.1 µl [³²P]dCTP (3000 Ci mmol⁻¹, 10 µCi µl⁻¹) (Amersham Nederland, 's-Hertogenbosch, The Netherlands) and 0.06 U of Super Taq DNA polymerase (Sphaero Q, HT Biotechnology, Cambridge, UK). Samples were covered with a drop of mineral oil (Sigma, St Louis, MO, USA) and were passed through a temperature cycling programme consisting of 4 min at 94°C, 33 cycles of 1 min at 94°C, 2 min at 55°C, 1 min at 72°C followed by an additional extension step of 6 min at 72°C in a thermocycling machine (MJ Research, Watertown, MA, USA). Samples were denatured by addition of a drop of 0.3% xylene cyanol, 0.3% bromophenol blue, 10 mmol l⁻¹ EDTA pH 0.90% (v/v) formamide and subjected to electrophoresis on a 0.4-mm-thick 6.5% polyacrylamide gel containing 7 mol l⁻¹ urea. Gels were dried and exposed to X-ray films (Konica, Tokyo, Japan) for periods ranging from 12 to 96 h. Results were interpreted by visual comparison of the allele intensities.

Table I

| Case | Histology     | Tumour a Location | Tumour a Percentage | Tumour b Location | Tumour b Percentage |
|------|---------------|-------------------|---------------------|-------------------|---------------------|
| OV1  | Clear cell    | Paraffin          | Left ovary          | 50%               | Right ovary         | 50%               |
| OV2  | Endometrioid  | Paraffin          | Left ovary          | 80%               | Right ovary         | 70%               |
| OV3  | Serous        | Paraffin          | Left ovary          | 70%               | Right ovary         | 80%               |
| OV4  | Undifferentiated | Paraffin       | Left ovary          | 90%               | Right ovary         | 90%               |
| OV5  | Serous        | Paraffin          | Left ovary          | 30%               | Right ovary         | 30%               |
| OV6  | Endometrioid  | Paraffin          | Left ovary          | 80%               | Right ovary         | 90%               |
| OV7  | Serous        | Paraffin          | Left ovary          | 10%               | Right ovary         | 50%               |
| OV8  | Endometrioid  | Paraffin          | Left ovary          | 50%               | Right ovary         | 60%               |
| OV9  | Serous        | Paraffin          | Left ovary          | 50%               | Right ovary         | 60%               |
| OV10 | Undifferentiated | Frozen          | Left ovary          | 70%               | Right ovary         | 50%               |
| OV11 | Serous        | Frozen            | Left ovary          | 20%               | Right ovary         | 30%               |
| OV12 | Undifferentiated | Frozen          | Left ovary          | 80%               | Right ovary         | 60%               |
| OV13 | Undifferentiated | Unavailable  | Left ovary          | 40%               | Right ovary         | 80%               |
| OV14 | Serous        | Frozen            | Right ovary         | 10%               | Left ovary          | 40%               |
| OV15 | Serous        | Frozen            | Left ovary          | 90%               | Right ovary         | 90%               |
| OV16 | Serous        | Frozen            | Left ovary          | 70%               | Right ovary         | 90%               |

*Sorted on the basis of DNA ploidy before DNA isolation.

Table II

| Marker | Chromosomal location | Heterozygosity |
|--------|----------------------|---------------|
| D1S162 | 1p32                | 0.91          |
| D1S175 | 1p21-q12            | 0.80          |
| APOA2  | 1q21-q23            | 0.74          |
| D1S158 | 1q32-q41            | 0.89          |
| D1S103 | 1q32-pter           | 0.88          |
| D3S1270| 3p                  | 0.75          |
| D3S11  | 3q21-q24            | 0.93          |
| GLUT2  | 3q26-q26.3          | 0.91          |
| D1S2323| 3q21                | 0.75          |
| IGF2R  | 6q25-q27            | 0.42          |
| F13A1  | 6q22-q24            | 0.78          |
| D6S89  | 6p24-p23            | 0.92          |
| D6S251 | 6q13-q21            | 0.78          |
| D6S249 | 6q13-q21            | 0.46          |
| D11S554| 1p12-p11.2          | 0.91          |
| D11S875| 1p12-p11.2          | 0.90          |
| D11S871| 1p12-p11.2          | 0.77          |
| TP5    | 17p13.1             | 0.68          |
| D17S515| 17q22-ter           | 0.88          |
| D17S520| 17q12               | 0.77          |
| D17S513| 17q13               | 0.89          |
| D17S579| 17q12-q21           | 0.87          |
| D17S250b| 17q11.2-q12        | 0.94          |
| D17S388| 17q22-23            | 0.68          |
| 46Es   | 17q23-24            | 0.74          |
| HOX2B  | 17q21.1-q21.3       | 0.82          |
| D18S40 | 18p11.21-ter        | 0.82          |
| D18S35 | 18q11-q12           | 0.72          |
| D18S54 | 18q11               | 0.81          |
| MBP    | 18q22-pter          | 0.80          |
| D22S156| 22q11               | 0.78          |
| IL-2RB | 22q11.2-q12         | 0.91          |
| CYP2D  | 22q13               | 0.80          |
| KALLMAN| Xp22.3              | 0.61          |
| DXS453 | Xp11.23-q21.1       | 0.72          |
| DXS454 | Xq21.1-q23          | 0.75          |

Statistics

The difference (d) between the multiple DIs can be caused by a measurement error or by a true biological difference. Previously Smit et al. (1990) derived a likelihood ratio (LR) that determines the probability that d is caused by matched pairing vs random pairing:

\[ LR = \frac{\sigma_2}{\sigma_1} e^{-\frac{d^2}{2\left(\frac{\sigma_1^2}{\sigma_2} + \frac{\sigma_2^2}{\sigma_1}\right)}} \]

Where \( \sigma_1 \) is the measurement error variance of the natural logarithm of the DI (LDI). The calculated value for \( \sigma_1 \) was 0.041, based on repeated flow cytometric experiments (Kute et al., 1988). \( \sigma_2 \) is the variance of LDI of randomly paired DIs of ovarian tumours and was determined to be 0.24 (Smit et al., 1990).
Statistical analysis of the LOH data was performed according to Jacobs et al. (1992) with minor modifications. Where $n$ deposits of a tumour have loss of heterozygosity for a polymorphism, the probability of each deposit losing the same allele as an independent even is $(1/2)^{n-1}$. This approach does not take the locus-specific chance for allele loss into consideration. In order to do this, chromosome arm-specific LOH frequencies were derived from the pooled data of four ovarian cancer allelotype studies (Sato et al., 1991; Cliby et al., 1993; Yang-Feng et al., 1993; Osborne and Leech, 1994).

For a locus $i$ the LOH frequency is $x_i$. If LOH of locus $i$ is observed for tumour $a$, the probability that LOH occurred at this locus in tumour $b$ independently is $x_i$, and the probability that the same allele is involved is $x_i$. The probability that in a specific case all observed concordant LOH patterns occurred as independent events is equal to the product of the single probabilities and is designated as the probability of independent origin (PIO). In the calculation of the PIOs we used a conservative approach. For example, when multiple markers on the same chromosomes showed LOH (indicating loss of a whole chromosome) only the highest LOH frequency was included in the PIO calculation since these multiple LOH patterns were probably not independent events.

The same was done for cases where identical chromosomal breakpoints were found. It should be considered however, that the probability that chromosomal breakpoints independently occur in the same region in two tumour deposits will probably be lower than the regional LOH frequency.

**Results**

**Flow cytometry**

Nuclear suspensions of all samples were prepared for DNA ploidy analysis. In eight of the 16 cases (OV1, OV2, OV3, OV4, OV6, OV7, OV15 and OV16) the DI was identical in both tumour sites investigated, in five cases (OV8, OV9, OV11, OV12 and OV14) the difference in DI was relatively small (0.1) and in three cases DIs between 0.2 and 0.9 were observed (OV5, OV10 and OV13) (Table III). In Figure 1 the DNA histograms of tumour a and tumour b from case OV14 are shown. Both histograms show an aneuploid population with almost similar DIs (1.7 and 1.6) with varying fractions of diploid cells. The LR of 1.39 calculated for this case is slightly in favour of a unifocal origin. In a total of 14 cases DIs were compatible with a unifocal origin (LR range 1.39–5.85). The extremely low LRs found for cases OV5, OV10 and OV13 made it unlikely that the differences in DI were caused by measurement errors, and thus indicate true biological differences.

**Detection of LOH**

Samples from 24 out of 32 tumours were enriched for their tumour cell content by flow cytometric cell sorting based on DNA aneuploidy (Table I). The effect of the enrichment is demonstrated in Figure 2. Both sorted diploid fractions of OV11 show two constitutional alleles for microsatellite marker D18S34, while a relative intensity decrease of the upper allele in the unsorted samples was observed, strongly suggesting LOH which is confirmed by PCR amplification of the aneuploid fraction showing a complete loss of the upper allele. The results of comparative LOH analysis of the 16 tumour pairs are summarised in Figure 3. Fourteen cases were informative concerning LOH. The cases were divided into three groups according to concordant or discordant LOH patterns. Group I includes 12 cases showing concordant LOH and ROH patterns for all informative markers. Within this group cases are ranked according to the number of loci showing concordant LOH, ranging from six (case OV13) to zero concordant LOH patterns. Case OV7 was also included in this group although no LOH was detected at any of the investigated loci. For case OV6 no signal was obtained for chromosome 22 in tumour b but concordance was observed for all other markers. The calculated probabilities that the observed LOH patterns could have occurred by coincidental loss of the same allele at all loci ranged from 0.000021 to 0.28 (Table III). Because of the absence of concordant LOH patterns no probabilities could be calculated for cases OV7 and OV8. The LOH data and the calculated probabilities are in most cases highly suggestive for a unifocal origin of the tumours.

Cases in group III show discordance for all observed LOH patterns. Loss of opposite alleles was not observed.
Remarkably, in both cases LOH was found in only one tumour site (OV3b and OV10b).

Group II consists of two cases which showed concordant as well as discordant LOH patterns. Figure 4 shows the partial allelotype for four informative microsatellite markers of case OV11. LOH can be observed in the aneuploid fraction of tumour a as well as in the aneuploid fraction of tumour b for D17S588 since in both lanes the upper allele is completely lost, while ROH was observed for marker D3S1270. However, only the aneuploid fraction of tumour a shows LOH for the chromosome 6 markers F13A1 and D6S251 while ROH for tumour b was observed. The markers for the other chromosomes showed concordant patterns (LOH for chromosomes 17, 18p and 22 and ROH for chromosomes 3, 11, 18q and X). Together with identical chromosome 18 breakpoints in both tumours (between D18S34 and D18S40) this strongly suggests a unifocal origin. This is supported by the calculated PIO of 0.0089. Although in case OV9 an independent origin cannot be excluded on the basis of a PIO of 0.28, the presence of identical chromosome

| Group | Chromosome |
|-------|------------|
|       | 1  3  6  11  17  18  22  X |
| Case  | ab ab ab ab ab ab ab ab |
| 1     | ab ab ab ab ab ab ab ab |
| 2     | ab ab ab ab ab ab ab ab |
| 13    | ab ab ab ab ab ab ab ab |
| 16    | ab ab ab ab ab ab ab ab |
| 12    | ab ab ab ab ab ab ab ab |
| 15    | ab ab ab ab ab ab ab ab |
| 1     | ab ab ab ab ab ab ab ab |
| 4     | ab ab ab ab ab ab ab ab |
| 6     | ab ab ab ab ab ab ab ab |
| 14    | ab ab ab ab ab ab ab ab |
| 5     | ab ab ab ab ab ab ab ab |
| 7     | ab ab ab ab ab ab ab ab |
| 8     | ab ab ab ab ab ab ab ab |
| 11    | ab ab ab ab ab ab ab ab |
| 9     | ab ab ab ab ab ab ab ab |
| 10    | ab ab ab ab ab ab ab ab |

Figure 3 Graphic representation of the allelotypes of the ovarian tumours for chromosomes 1, 3, 6, 11, 17, 18, 22 and X. The left chromosome of each chromosome pair corresponds to tumour a and the right chromosome to tumour b of each case. The small numbers indicate the number of informative markers for each chromosome. Black symbols indicate LOH, white symbols ROH. When microsatellite markers could not properly be amplified or where all tested markers were homozygous for a specific chromosome no chromosome symbol was drawn. Case numbers are indicated at the left side. The cases are grouped according to the occurrence of concordant LOH patterns. Within the groups the cases were ranked according to descending number of observed LOH events.
17 breakpoints, identified by four informative markers is still strongly suggestive of a unifocal origin of both tumours.

**LOH vs DNA index**

In group I identical allelotypes correlate to equal or nearly equal DIs (d = <0.1) in all cases except OV5 and OV13. However, the P10 values show a much higher variation than the LRIs for these cases. Although, for OV13 the LOH patterns are totally identical for all informative markers on seven different chromosomes (PIO = 0.000021), a large difference in DI was observed (LR = 1.29 x 10⁻⁸). This discrepancy can be explained by assuming that the aneuploid stemline in tumour a represents a tetraploidised subclone of the diploid stemline of tumour b. This subclone must have originated after the establishment of the observed allelotype. In case OV5 neither DNA ploidy analysis nor the allelotype can exclude an independent origin. Although both tumours lost the complete chromosome 17 this still could be an independent event since loss of chromosome 17 is very common in ovarian tumours. Despite a non-complete identity of the allelotypes of the two cases in group II the FCM results are also in favour of a unifocal origin of the tumours. In group III the discordant LOH patterns correlate with differences in DNA index for case OV10 while the two methods gave contradictory results for OV3.

**Discussion**

In the present study two different genetic approaches were used to study the origin of bilateral ovarian tumours. We found evidence for a unifocal origin in 14 of 16 cases on the basis of LOH analysis, whereas DNA ploidy analysis did not always provide conclusive evidence about the origin of tumours. The use of DNA ploidy analysis is based on the assumption that identical, aneuploid DIs are not likely to be the product of an independent ploidy evolution but rather reflect a unifocal origin. DNA FCM is a rapid technique which is used in many clinical pathological laboratories throughout the world. A limitation to this type of analysis is that ploidy evolution may continue after the formation of metastases, e.g. by tetraploidisation. This would result in a DI difference that might be interpreted as evidence against a unifocal origin. An illustration of this phenomenon is case OV13 in which DIs of 1.9 and 1.0 were observed while the allelotype was completely identical including LOH at six different chromosomes (Figure 3). Tetraploidisation apparently occurred after establishment of the allelotype. Also in the case of DI heterogeneity in the primary tumour metastasis of a minor subpopulation not detected in the primary tumour might yield a similar DI difference. Apart from this the limited resolution of DNA FCM which on average cannot detect DNA content differences less than 5% may lead to spurious identity of DIs.

Microsatellite-based LOH analysis can detect genetic differences at the subchromosomal level and is intrinsically more sensitive and specific. In contrast to DNA ploidy analysis, ongoing clonal evolution at different tumour locations might not preclude the identification of a unifocal origin owing to the clonal retention of acquired genetic aberrations, e.g. the two cases OV9 and OV11 in group II. The specificity of this approach depends on the number of independent LOH events as well on the a priori probability that a certain chromosome region will show LOH, e.g. regions harbouring tumour-suppressor genes. For instance, a coincidental, identical LOH pattern for chromosome 17 will be more likely than for chromosome 1. We have corrected for this in our calculations. With the present availability of highly polymorphic micro-satellite markers, the accurate mapping of chromosomal breakpoints can substantially increase the power of this type of analysis and provide definite answers about the clonality of multiple tumours. Identical chromosomal breakpoints are a strong indication of a unifocal origin. The accuracy by which the exact position of breakpoints and thus their identity can be determined, depends on the proximity of the flanking microsatellite markers. Other PCR-based methods that can provide information about the origin of multiple tumours are determination of X-chromosome inactivation (Allen et al., 1992), oncogene amplification (Li et al., 1994) and determination of oncogene or tumour-suppressor gene mutations. The use of X chromosome inactivation is hampered by the fact that half of the multifocal tumours will have identical inactivation just by chance. Oncogene amplification is less informative because (1) a limited number of chromosomal regions is known to be involved and (2) certain amplicons are strongly correlated with certain tumour types which makes an independent amplification in both tumours rather likely and therefore less appropriate. Mutation analysis of frequently involved oncogenes or tumour-suppressor genes like p53 is highly informative and applicable in many kinds of tumours including ovarian tumours. Disadvantages are the lack of mutations in a high percentage of tumours (for ovarian tumours...
about 50%), presence of mutational hot spots (ras, p53) and the possibility that a mutation occurs after metastatic spread.

A limitation of LOH analysis on DNA extracted from total tumour tissue is the fraction contaminating non-neoplastic cells. Under optimal conditions the minimum amount of tumour cells should exceed 40% (Gruis et al., 1993). We recently showed that tumour specimens with lower amounts of neoplastic cells can be made accessible for PCR-based LOH analysis by flow sorting of tumour cells on the basis of DNA ploidy and marker expression (Abeln et al., 1994). In the present study this enabled us to successfully enrich 24 of the 32 tumours where otherwise 18 tumours would have been excluded owing to low tumour cellularity.

In 10 of the 16 investigated cases statistical evidence of unifocal origin was found on the basis of LOH and in two cases (OV5 and OV9) evidence was obtained which suggested unifocal origin but failed to reach the level of statistical significance. For case OV9 the identical chromosome 17 breakpoint in both tumours provides strong evidence for a common origin of the different tumour sites. These findings are in agreement with previous reports based on DNA flow cytometry, karyotyping, p53 mutation analysis or allelotyping (Smitt et al., 1990; Pevovic et al., 1991; Jacobs et al., 1992; Mok et al., 1992) in which 31, 43, 11, 9 and 15 17 resistant events were observed.

From a theoretical point of view formal proof that tumours are multifocal is quite difficult. Only if the initial molecular event is known (as in e.g. APC mutation in colorectal adenoma development; Powell et al., 1992) discordance for this marker will be direct proof of multifocality.

The tumours in group III showed discordant LOH which suggests but does not prove multifocality since post-metastatic LOH does occur as evidenced by the tumours in group II.

The PLO calculation is correct under the condition that LOH is non-parental origin-specific. If a certain chromosomal region harbouring a tumour-suppressor gene is imprinted, loss of the other allele may totally switch off its suppressor function and will provide selective advantage with regard to carcinogenesis. The chance that two tumours in a patient independently lose the non-imprinted allele therefore may be higher than in a non-imprinted situation. In this case the PLO would be a less accurate measure for independent origin. The same holds for cases where LOH is the second hit uncovering a germline mutation according to Knudson's two-hit tumour-suppressor gene model, but this will be restricted to familial tumours. Parental specific LOH has been reported for some cases (reviewed by Feinberg, 1993) concerning Wilms' tumour, rhabdomyosarcoma, bilateral retinoblastoma, acute myelogenous leukaemia and neuroblastoma concerning regions on chromosomes 1, 2, 7, 11 and 13. However this phenomenon has not been reported for ovarian cancer. Although LOH at chromosome 11 is a frequent event. However both for the imprinted as well as in familial tumours, partial LOH generating a chromosomal breakpoint, would still enable discrimination of multifocal from unifocal origin. In our series chromosomes 6, 11, 17 and 18 were most frequently involved in LOH. LOH at 1q was previously reported to be a common event (51%) in ovarian tumours (Saito et al., 1992), although a candidate tumour-suppressor gene has not yet been reported in this region. WT1 may be a target gene for the loss of chromosome 11 (Eccles et al., 1992; Foulkes et al., 1993). TP53 could be the target gene for the chromosome 17p loss, since the gene is frequently involved in ovarian cancer (Milner et al., 1993). For BRCAl which also maps to chromosome 17, recently both germline (Futreal et al., 1994) as well as somatic mutations have been found in ovarian cancer (Hosking et al., 1995; Merajver et al., 1995) although a second gene on 17q may be a more likely target (Saito et al., 1993). Frequent LOH at 18q (60%) has also previously been reported, however the smallest region of overlap seems to exclude the DCC tumour-suppressor gene (Chenevix-Trench et al., 1992).

Interestingly two cases, OV9 and OV11, showed discordant LOH patterns for chromosomes 6 and 18 and concordance was observed at other loci. In contrast, relatively high LOH frequencies for chromosomes 6 (62%) and 18 (43%) were reported by Cliby et al. (1993). Our results suggest that LOH at chromosomes 6 and 18 may represent late events in tumour progression which in some tumours may not be essential for establishing the malignant phenotype. Whether the absence of LOH at chromosomes 6 and 18 respectively, in one of the sites definitely identifies these as the primary tumours is still unclear. Evidence for clonal divergence was previously reported by Jacobs et al. (1992) although in this study fewer loci were investigated.

Our results support the evidence for a monoclonal origin of the majority of bilateral ovarian carcinomas. PCR-based LOH analysis proved to be a more reliable technique for clonality determination than DNA FCM. The results obtained with both techniques indicate that clonal divergence after metastasis is an infrequent phenomenon in ovarian cancer.

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