Chromosome 6p amplification and cancer progression

Gda C Santos, M Zielenska, M Prasad, J A Squire

Chromosomal imbalances represent an important mechanism in cancer progression. A clear association between DNA copy-number aberrations and prognosis has been found in a variety of tumours. Comparative genomic hybridisation studies have detected copy-number increases affecting chromosome 6p in several types of cancer. A systematic analysis of large tumour cohorts is required to identify genomic imbalances of 6p that correlate with a distinct clinical feature of disease progression. Recent findings suggest that a central part of the short arm of chromosome 6p harbours one or more oncogenes directly involved in tumour progression. Gains at 6p have been associated with advanced or metastatic disease, poor prognosis, venous invasion in bladder, colorectal, ovarian and hepatocellular carcinomas. Copy number gains of 6p DNA have been described in a series of patients who presented initially with follicle centre lymphoma, which subsequently transformed to diffuse large B cell lymphoma. Melanoma cytogenetics has consistently identified aberrations of chromosome 6, and a correlation with lower overall survival has been described. Most of the changes observed in tumours to date map to the 6p21–p23 region, which encompasses approximately half of the genes on all of chromosome 6 and one third of the number of CpG islands in this chromosome. Analyses of the genes that cluster to the commonly amplified regions of chromosome 6p have helped to identify a small number of molecular pathways that become deregulated during tumour progression in diverse tumour types. Such pathways offer promise for new treatments in the future.

Abnormalities of chromosome number leading to aneuploidy are the most common cytogenetic aberrations in cancer. Both genomic and cytogenetic analyses of different types of tumours have shown recurrent DNA copy-number increases associated with progression. A clear association between DNA copy-number aberrations and prognosis has been found in a variety of tumours.1

Partial or whole gain of 6p is much more common than loss. Molecular cytogenetic analysis of chromosome 6p has shown three modes of 6p gain:

1. The most frequent mechanism for generating 6p gain is the formation of an isochromosome, in which centromeric misdivision leads to triplication of 6p and monosomy of 6q;
2. Acquisition of an unbalanced chromosome translocation can also lead to partial or complete 6p gain; and
3. Focal amplification of a discrete region of 6p typically involves a much larger copy number increase (>5-fold).

The last mechanism of gain may involve the formation of double minute chromosomes or homogeneous staining regions.

Genomic imbalance can be detected and mapped by comparative genomic hybridisation (CGH). This method has been used to detect copy-number increases on chromosome 6p in several human neoplasias,2–7 suggesting that this region may contain genes involved directly or indirectly in the pathogenesis and pathways of cancers. This review focuses on gains and amplifications that have been established to be recurrent for chromosome 6p in different classes of solid tumours (fig 1 and table 1), with emphasis on the most commonly amplified genomic interval of 6p21–p23. Such observations are strongly suggestive that this central part of the short arm of chromosome 6p harbours one or more oncogenes directly involved in tumour progression.

CARCINOMAS

In carcinomas, there is a general association between gains of 6p and tumour progression. In bladder carcinomas, gains of 6p have been reported in invasive bladder tumours, present in 7–55% of patients.8–11 Significant associations have been found between 6p22 gain and high histological grade,12,13 high tumour cell proliferative activity14 and metastases at initial presentation,15 suggesting that acquisition of 6p gain may confer a growth advantage and lead to disease progression.

As in advanced stages of transitional cell carcinomas in vivo, gains and amplifications of 6p were often found in transitional cell carcinoma lines with a common region of amplification at the 6p21.3–p23 locus.17 Over-representation of 6p22 was the only individual change that was significantly linked to high tumour grade in a series of 54 pT1 urinary bladder carcinomas. The risk of progression was significantly associated with the number of deletions, but not with the number of gains.18 Moreover, gain at chromosome 6p22–p23 was noted in two patients with metastasis at diagnosis among pT4a, pT1 and pT2–T4G3 tumours and gains of 6p and 10p were more frequent in pT1G3 compared to pT1G2 tumours.19 In high-stage lesions 6p22 gains occurred as late events,20 and this gain was the only CGH change strongly associated to a high proliferative activity in the

Abbreviations: BCC, basal cell carcinoma; CGH, comparative genomic hybridisation; CRC, colorectal carcinoma; HCC, hepatocellular carcinoma

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Figure 1 A map of the 6p21–p23 cytobands showing regions of gains and amplifications for specific tumours with genes that were previously published. The black up arrow indicates gain/amplification to the pter and the black down arrow indicates gain/amplification to the centromere. The solid black bar indicates the minimal region of gain; the white bar with black hashed lines indicates a region of amplification; the white bar with black vertical lines indicates a focal gain; and the back bar with white hashed lines indicates a region of gain for the whole 6p arm.

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tumour cells and independent of grade and stage of the tumour.14

In small-cell carcinomas, which represent a rare histological subtype of urinary bladder cancer, gains of DNA sequences were most prevalent at 8q, 5p, 6p and 20q.29 High-level amplifications were detected most often at 6p22.3 (E2F3) and at four other genomic locations in 41 primary bladder tumours. Interestingly, there was a significant complementary association between gain of cyclin D1 at 11q13 and gain of E2F3, although there was no significant relationship between copy number changes and tumour stage or grade.30 Using quantitative multiplex polymerase chain reaction to study DNA from 59 bladder tumours, the focal region of genomic gain on 6p22 was mapped to a minimal region, spanning a genomic distance of 0.5 Mb.15 The E2F3 gene has been recently implicated as the target of 6p22 genomic gain in bladder cancer.22, 23 In a study using bladder tumour-derived cell lines, NM_017774 showed an expression related to the 6p22.3 amplicon.24 A recent paper showed overexpression of ID4 gene, which maps between E2F3 and DEK, in bladder cancer.25

Increased copy number of chromosome arm 6p has been associated with advanced stages of colorectal cancer (Dukes’ stage D) and metastasis.7 In a recent CGH study on liver metastasis of colorectal carcinoma (CRC), gain in 6p21 was found in 41% of the patients. Taken together, both these observations suggest that acquisition of 6p gain may be contributing to progression in colorectal cancer, and that part of the short arm of this chromosome might harbour one or more oncogenes.26 A subtractive CGH analysis using paired samples from 20 patients with CRC with primary tumours and synchronous or metachronous liver metastases detected frequent gains in DNA copy number at 6p. Analysis of 11 CRC cell lines using array-based CGH showed one 6p candidate gene, cyclin D3, which was significantly upregulated by quantitative reverse transcriptase-polymerase chain reaction in liver-metastatic lesions compared with primary lesions.

Table 1 Tumours with 6p21–p23 gain or amplification

<table>
<thead>
<tr>
<th>Type of tumour</th>
<th>Number of tumours</th>
<th>Tumours with gain in 6p21–p23 (%)</th>
<th>Tumours with amplification in 6p21–p23 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carcinomas*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatocellular carcinoma</td>
<td>409</td>
<td>22.61</td>
<td>0.20</td>
</tr>
<tr>
<td>Merkel cell carcinoma</td>
<td>48</td>
<td>27.10</td>
<td>2.10</td>
</tr>
<tr>
<td>Basal cell carcinoma</td>
<td>16</td>
<td>40.30</td>
<td>0.00</td>
</tr>
<tr>
<td>Ovarian serous carcinoma</td>
<td>56</td>
<td>28.60</td>
<td>0.00</td>
</tr>
<tr>
<td>Transitional cell carcinoma</td>
<td>133</td>
<td>9.68</td>
<td>0.51</td>
</tr>
<tr>
<td>Lymphoid tumours</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Large B cell lymphoma</td>
<td>360</td>
<td>8.36</td>
<td>0.07</td>
</tr>
<tr>
<td>Plasmacytoma</td>
<td>21</td>
<td>22.76</td>
<td>0.00</td>
</tr>
<tr>
<td>Sarcomas</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Osteosarcoma</td>
<td>137</td>
<td>33.19</td>
<td>3.39</td>
</tr>
<tr>
<td>Malignant peripheral nerve sheath</td>
<td>70</td>
<td>23.93</td>
<td>0.00</td>
</tr>
<tr>
<td>Leiomyosarcoma</td>
<td>136</td>
<td>9.98</td>
<td>1.64</td>
</tr>
<tr>
<td>Melanomas</td>
<td>91</td>
<td>31.78</td>
<td>0.00</td>
</tr>
<tr>
<td>Retinoblastoma</td>
<td>133</td>
<td>39.36</td>
<td>10.50</td>
</tr>
<tr>
<td>Glioblastoma</td>
<td>108</td>
<td>4.60</td>
<td>0.90</td>
</tr>
<tr>
<td>Neuroblastoma</td>
<td>303</td>
<td>22.10</td>
<td>0.17</td>
</tr>
<tr>
<td>Carcinomas</td>
<td>23</td>
<td>21.70</td>
<td>6.31</td>
</tr>
</tbody>
</table>

*This site does not currently allow the determination of the level of imbalance in a subset of breast and colon carcinomas.

Summary of genomic imbalance datasets from Progenetix (http://www.progenetix.com).

CGH studies have shown that the most frequent gains in hepatocellular carcinoma (HCC) occurred at 8q, 1q, 3q, 6p and 17q.31–33 Chromosomal aberrations investigated in HCC cell lines that had hepatitis B virus integration showed chromosomal gains at 6p.34 Amplifications at 1q and 6p appeared to be independent factors for venous invasion in HCC.35 Significantly, a recent study described that physical clusters of two or more genes within predefined distance thresholds were detected non-randomly on chromosomal regions 1q, 6p, 8q, 20q and Xq, indicating that HCC-related genes are physically clustered at specific chromosomal locations.36 A previous study that mapped genes overexpressed in HCC to chromosomal locations found 13 regions of frequent cytogenetic change, including 6p gain.37 A recent explorative CGH meta-analysis of HCC showed that the most marked amplifications were present at 1q (57.1%), 8q (46.6%), 6p (22.3%) and 17q (22.2%).38

In breast carcinomas, genomic imbalances with statistical significance were observed in poorly differentiated (G3) and oestrogen receptor negative tumours, including one region of 6p gain.39 One of the most frequent chromosomal aberration found in patients with high-risk stage II/III breast cancer was 6p gain.40 Gain on 6p21 was identified in 31 (45%) formalin-fixed and paraffin-wax-embedded primary advanced breast tumours analysed by microarray-based CGH.

Recurrent gains of chromosome 6 have been detected in Merkel cell carcinomas.41 Although no significant correlation between genomic aberrations and clinicopathological factors was shown, primary tumours expressing changes in DNA were predominantly distinguished in large Merkel cell carcinomas, and risk of metastatic dissemination was threefold compared with tumours without chromosomal changes.42

In basal cell carcinoma (BCC) of the skin, recurrent changes were observed in 47% of the patients at 6p21-pter. Interestingly, it was found that regional gain of 6p was strongly associated with 6p gain; however, no correlation between this association and clinical or histological appearance was observed in these patients.7 These results correlate with a previous study that identified clonal trisomy of chromosome 6 in BCC direct preparations by conventional cytogenetics and fluorescent in

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situ hybridisation analysis. No correlations between the BCC cytogenetic results and clinical parameters such as site, age, sex and recurrence rate have been shown. 69

Gains of 6p were more common in stage IIIc than in stage IIIa+ b ovarian serous papillary adenosarcomas. 70 A study of ovarian epithelial tumours using CGH found that high-grade serous carcinomas had more than twice as much chromosomal imbalance as low-grade serous carcinomas and also had pronounced changes. Overlapping changes occurring in serous and non-serous carcinomas were gains on 3q and 6p, as well as losses on 4q. Among other chromosomal imbalances, 6p gain was associated with poor prognosis of ovarian carcinomas, 4 and 6p22.1–p21.2 was one of several regions associated with acquisition to drug resistance in serous carcinomas. 42

LYMPHOID TUMOURS

Amplification of the 6p21 locus has been documented in non-Hodgkin’s lymphoma. Among 27 specimens of patients with plasmacytoma, 25% of the specimens showed gain of 6p, with a minimal overlapping region at 6p21.3-pter. 71 Acquisition of 6p has a significantly higher frequency in large B cell tumours, and it was an independent prognostic factor with losses of 11q21–q23.1 and 17p. 72 In another study of diffuse large B cell lymphoma with primary tumours and recurrent tumours excised after chemotherapy or radiotherapy, 21% showed 6p gain with minimal common regions at 6p22-pter. High-level amplifications were observed at 6p23-pter and occurred only in different recurrent tumours. 73 In a series of 23 patients, 21 showed genomic changes, with a mean value of 11 aberrations/sample (range 2–29). Among other regions, the minimal common region of 6p22-pter was most frequently gained. 74 In non-Hodgkin’s lymphomas, it has been shown and suggested that gains on 6p among other changes may be important in the transformation from low-grade to high-grade disease. 75 Recently, over-representation of 6p12.3–p21 among other genomic aberrations was reported in samples from patients who presented initially with follicle centre lymphoma, which subsequently transformed to diffuse large B cell lymphoma as measured by array CGH. 76 A subset of genes that mapped to locations of gain or amplification exhibited coordinated increase or decrease of expression. 76

MELANOMAS

Melanoma cytogenetics has consistently identified aberrations of chromosome 6 as the most common change. CGH analyses have identified recurrent chromosomal abnormalities at several locations including 6p gain in choroid and ciliary body melanomas (30–85%), 76–78 in acral and superficial, 79–80 and in sinonasal mucosal melanomas (93%). Gain of 6p occurred frequently in combination with a loss of copy number or a normal copy number of the opposite chromosome arm, suggesting isochromosome formation. 77

Changes on 6p, detected by microsatellite analysis in 15 of 30 (50%) patients with uveal melanoma, were preferentially observed in tumours without chromosome 3 changes. 78 Another study of 12 patients with uveal melanomas showed under-representation of chromosome 3 exclusively in tumours with ciliary body involvement, whereas anomalies of chromosome 6 were described only in pure choroidal melanomas. 80 Recently, an inverse relationship between monosomy 3 and gains on 6p, detected by microsatellite analysis in 15 of 30 (50%) patients with uveal melanoma, were preferentially observed in tumours without chromosome 3 changes. 78 Another study of 12 patients with uveal melanomas showed under-representation of chromosome 3 exclusively in tumours with ciliary body involvement, whereas anomalies of chromosome 6 were described only in pure choroidal melanomas. 80 Recently, an inverse relationship between monosomy 3 and gains on 6p, detected by microsatellite analysis in 15 of 30 (50%) patients with uveal melanoma, were preferentially observed in tumours without chromosome 3 changes. 78 Another study of 12 patients with uveal melanomas showed under-representation of chromosome 3 exclusively in tumours with ciliary body involvement, whereas anomalies of chromosome 6 were described only in pure choroidal melanomas. 80

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RETINOBLASTOMA

Gains of the entire chromosome arm 6p have been reported very frequently in retinoblastoma. 2, 74–77 Interestingly, by far the most frequent mechanism of gain is by acquisition of an isochromosome of 6p. 2, 74–76, 80, 82 Karotype studies showed small additional fragments of 6p, often including the distal segment 6p25–p22. This led to the assumption that increased copy number of genes localised in this region of 6p may confer a selective growth advantage to retinoblastoma cells. 81 One of the most frequent imbalance observed in a series of primary retinoblastoma (13/24) cases was 6p gain, and no evident selection was found between any of the imbalances identified and either the differentiation or the histopathologic risk. 83 Gains at 6p were also identified in another CGH study in 11 (42.3%) tumours, suggesting that 6p22–p25 might be a candidate chromosome region for containing presumptive retinoblastoma growth-promoting genes. 84 Subsequent CGH results have shown that the common region of gain maps to chromosome band 6p22.4 Using quantitative multiplex polymerase chain reaction, the same group narrowed down the region to a 0.6-Mb region spanning the UniSTS markers X64229 and WI-19208. 85 Recently, it was found that only three of seven genes, DEK, NUP153, and EZF3, in this region analysed using real-time reverse transcriptase-polymerase chain reaction are overexpressed in retinoblastoma tumours and cell lines with comparison to those without gain, thus implying that acquisition of 6p or 1q gain may confer prognostic differences. 35
documented genomic gain of 6p22. E2F3 and DEK mRNA overexpression was always associated with protein overexpression of primary tumours relative to the adjacent normal retina. E2F3 was strongly expressed in actively proliferating cells, whereas DEK was overexpressed in all tumour cells. It was concluded that both E2F3 and DEK were promising targets of the 6p22 genomic gain in retinoblastoma.\(^8\) Another study narrowed the minimal region of gain on 6p to the band 6p22.3 using a panel of retinoblastomas and retinoblastoma cell lines. The data indicated that the minimal region of chromosomal imbalance in 6p22.3 includes the NUP153, KIF13A, TPMT, DEK, ID4, E2F3 and TTRAP genes. Copy number gains were most consistently identified in the region containing the DEK and E2F3, thus suggesting these genes as potential targets of 6p gains.\(^8\) A high-resolution matrix-CGH analysis performed on a series of 17 primary retinoblastomas and 4 retinoblastoma cell lines showed on 6p a minimally overlapping gained region at 6p21.31–p21.33 that was 2.31 Mb in size and suggested the tumour necrosis factor \(\alpha\) as a potential candidate gene.\(^7\)

### OTHER TUMOURS

One frequent over-representation was seen at 6p (minimal region of overlap in six patients, 6p22–p23) in a CGH analysis of 19 uterine and extrauterine carcinosarcomas. Amplifications of 6p were detected in two patients.\(^5\) Amplification and over-expression of cyclin D3 gene located in the 6p22 region has been detected in glioblastoma cell lines.\(^4\) It was suggested that the increased amounts of cyclin D3 caused by gene amplification could be involved in the development or progression of that glioblastoma.\(^4\) Common gains observed in 29 neuroblastomas were on chromosomes 7, 6, and 18. High-level amplifications were detected at 2p23–p25 (MYCN region), at 4q33–q35 and at 6p11–p22.\(^7\) A CGH analysis of clinically non-progressing stage 4s neuroblastomas showed a high rate of whole-chromosome aberrations (73.4%), with an over-representation of chromosomes 2, 6, 7, 12, 13, 17 and 18 and an under-representation of chromosomes 3, 4, 11 and 14.\(^8\)

### CONCLUSIONS AND FUTURE PROSPECTS

CGH studies have identified frequent amplifications and rearrangements involving 6p in a wide variety of different types of tumours. These observations suggest that genes more generally associated with tumour progression may map to this genomic location. In all, 624 genes are seen in the 33-Mb region of 6p21–23, which represents about half of the genes on whole chromosome 6 (table 2). Several lines of evidence presented in this review implicate that one or more putative oncogenes may underlie the association between gain of 6p and tumour progression. The genomic architecture of the 6p region is noteworthy. The number of CpG islands in the 6p21–p23 region corresponds to one third of the number of CpG islands in chromosome 6. Recently, it has been shown that gene density, guanine-cytosine content and replication timing all co-correlate strongly with transcriptional activity, thus confirming that expressed genes tend to be replicated early in the S phase.\(^9\)

In solid tumours, significantly high levels of chromosome abnormalities have been detected, but distinction between critical and consequential events has been a major challenge. It has been proposed that correlations between imbalances would show possible cytogenetic pathways. Some tumours described in this review present concordant genomic changes associated with 6p gain. In the future, the application of systematic higher resolution array CGH studies will help to show correlations and linked behaviour among genes that cluster to chromosome 6p and other specific regions, thus unravelling different pathways for an improved understanding of tumour progression. Analysis of the genes that cluster to the commonly amplified regions of chromosome 6p has helped to identify a small number of molecular pathways that become deregulated during tumour progression in diverse tumour types.\(^10\) Identification of such pathways will offer promise for new approaches to treatment in the future.

### Take-home messages

- A clear association between DNA copy-number aberrations and prognosis has been found in a variety of tumours. CGH studies have identified frequent amplifications and rearrangements involving 6p in different types of tumours.
- Several lines of evidence presented in this review suggest that a part of the short arm of chromosome 6p harbours one or more oncogenes directly involved in tumor progression.
- Some tumours described in this review present concordant genomic changes associated with 6p gain showing possible cytogenetic pathways. The application of systematic array CGH studies can help to show correlations and linked behaviour among genes that cluster to chromosome 6p and other specific regions. Such an approach will unravel different pathways for an improved understanding of tumour progression.

### Table 2 Genomic features of 6p21–p23

<table>
<thead>
<tr>
<th>Genomic features of:</th>
<th>6p21–p23 region</th>
<th>Chromosome 6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Size of the region (Mb)</td>
<td>33</td>
<td>171</td>
</tr>
<tr>
<td>Number of genes</td>
<td>624</td>
<td>1394</td>
</tr>
<tr>
<td>Repeats</td>
<td>59 854</td>
<td>265 300</td>
</tr>
<tr>
<td>CpG island</td>
<td>5508</td>
<td>17 622</td>
</tr>
<tr>
<td>Variations (SNPs)</td>
<td>141 464</td>
<td>601 204</td>
</tr>
<tr>
<td>OMIM morbid</td>
<td>62</td>
<td>121</td>
</tr>
<tr>
<td>Miltenberg breakpoint elements</td>
<td>78</td>
<td>665</td>
</tr>
</tbody>
</table>

SNP, single nucleotide polymorphism.


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