Gender and ploidy in cancer survival

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Abstract

Background Females carry a better prognosis than men for many cancer types. We hypothesized that chromosomal changes, in particular numerical alterations of the sex chromosomes or the presence of near-triploidy may contribute to these gender differences.

Methods To characterize the influence of gender a literature search was performed for survival data of 27 tumor types. All entities were categorized by the strength of evidence for differences in survival between females and males. To test our hypothesis the Mitelman database of chromosomal alterations was evaluated for the major tumor types occurring in both women and men. Numerical gonosome alterations were documented and mean chromosome numbers were converted into histograms to provide insight into the ploidy level of 37 cancer types.

Results In general, a survival advantage of women could be shown for most, but not all cancer types. In addition, 36,859 karyograms were analyzed. Numerical gonosome alterations were more frequent in males than females indicating a potential link with gender differences in survival. Neartriploidy was a common phenomenon in many cancer types suggesting that it represents a metastable condition of the cancer genome. It was not related to gender differences in survival. However, the extent of triploidy and aneuploidy was associated with poor prognosis in carcinomas. There was no single case in the Mitelman database with normal chromosome number (n=46) that did not carry at least one structural or numerical aberration.

Conclusions Our study highlights the importance of chromosomal changes in tumor formation and progression. In addition, it suggests potential associations with gender specific differences in survival.

Keywords Gender · Aneuploidy · Chromosomal chaos · Prognosis

1 Introduction

Observations showed that chromosomal alterations and aneuploidy are associated with the survival of tumor patients. Furthermore, it is known that there is an influence of sex on tumor survival. In this work, on the one hand, we wanted to systematically investigate whether and if so in which tumor types gender has an influence on the prognosis. As a predictive value we used the 5-year survival rate to compare the difference in survival between men and women. On the other hand the influence of ploidy on tumor survival should be investigated.

The term ploidy is related to the chromosome complement, but also to the DNA content of cells. Aneuploidy is the state of a cell in which there are numerical aberrations of chromosomes compared to normal. Aneuploidy may also occur when structural chromosomal aberrations are associated with the gain or loss of genetic material. These genetic imbalances are often detected by a change in the DNA content of cells [29]. Overall, aneuploidy is a common and essential characteristic of malignant tumors [20]. Some authors relate the state of aneuploidy exclusively to
numerical chromosomal aberrations while others include structural aberrations that are associated with the loss or gain of chromosomal material [16].

In this paper we used the term aneuploidy primarily in the context of numerical chromosomal aberrations. A correlation between aneuploidy and in particular the extent of chromosomal changes with the survival of cancer patients has already been shown long time ago [22]. However, in most studies only one tumor was examined for aneuploidy. There are only few studies which analyzed several tumor entities [5], [72].

Malignant tumors frequently exhibit DNA distributions in the near triploid range as we recently pointed out for lung cancer [60]. Triploidy in cancer can be regarded as evidence for aneuploidy. It is yet unclear whether this state of triploidy is related to specific biological or clinicopathological tumor characteristics.

Specifically, we asked ourselves in this work the following questions: 1) Which tumor entities show gender differences in survival? 2) Which tumor types harbour aneuploid chromosome numbers and have a near triploid chromosome set? 3) Is there a correlation between triploidy/aneuploidy and survival? 4) Is there a correlation between the ploidy level/triploidy or sex chromosome aberrations and gender-specific difference in cancer survival? More than 30 tumor entities were analyzed regarding their survival data and ploidy state.

2 Materials and methods

2.1 Gender-specific prognosis

In this work we examined whether gender could have an influence on the prognosis. As a predictive value, we used the 5-year survival rate to compare the difference in survival between men and women. We examined the overall survival of 34 tumor entities, including seven gender-specific tumor types. Data on gender-specific prognosis were retrieved from PubMed. We took into account preferentially studies in which the entire tumor entity was represented, i.e. no studies in which only certain stages or tumor subgroups (age, etc.) were considered.

In this respect, there are only few sources that reported comprehensively on multiple tumor entities. Our main sources of the survival rate were the Robert Koch Institute (RKI) and the Eurocare-3 study, this data mainly covered European and German populations.

In addition, many studies on individual tumor entities were retrieved in PubMed using the following search parameters: tumor entity, such as “lung cancer”, “epidemiology”, “sex specific survival” or “gender-specific prognosis”. The studies were then summarized in a table in which the evidence for gender-specific difference in prognosis is reflected by the number of studies reporting such differences (Table 1).

2.2 Tumor ploidy

To search for the ploidy in various tumors, we used the Mitelman database [44]. It represents only modal cancer cell lines and not all existing sidelines that are present in these tumors. With the option “Full Searcher” specific tumor types can be selected. The result is a table that lists the complete karyotypes of the different tumor cases. These karyotypes were transferred to Excel. For each karyotype, we recorded the total chromosome numbers. In some tumors, only a range of chromosome numbers were available (e.g. 64–70). For these cases a mean value was calculated, i.e. minimum plus maximum chromosome number divided by two, to get a single value for each case (i.e. 67 in the above mentioned example). The range of the chromosome numbers of these cases are available as supplementary data. Importantly, only three cases had numbers that varied between 46 (minimum) and 92 (maximum) chromosomes (1 colon carcinoma, 1 multiple myeloma, 1 thyroid carcinoma). These cases were eliminated from the ensuing analysis.

The numbers were then presented in the form of a histogram (Fig. 1). To simplify the representation, the tumors were grouped according to a range including cases with three adjacent chromosome numbers. That means for example that within the group of diploid tumors not only those cases with 46 chromosomes but also those with 45 and 47 were included. In general, the histograms represent the distribution of chromosome numbers within a tumor entity.

In order to define distinct ploidy states, the proportions of cases with hyperdiploid, predominant triploid, predominant hypertriploid and aneuploid sets of chromosomes were calculated. “Hyperdiploid” tumors were classified as those with chromosome numbers larger than 47 \( (n>47) \). Accordingly, cases with “predominant triploid” karyotypes were defined as tumors with chromosome numbers from 57 to 83 \( (57<n<83) \). “Predominantly hypertriploid” tumors were those with \( n>57 \). “Aneuploid” tumors were classified as those with chromosome numbers not in the diploid or tetraploid range \( (n\neq45–47 \text{ and } 90–92) \).

Furthermore, we calculated the proportion of cases with mutated sex chromosomes in different tumor entities, i.e. those who did not have the normal XX or XY karyotype. The comparison between the proportion of mutant sex chromosomes in tumor entities of women and men was then represented as a curve diagram (Fig. 3).
2.3 Survival analysis and statistics

The data from the ploidy analysis and the calculated proportion of mutated sex karyotypes were then correlated with the 5-year survival rates by calculating regression/correlation coefficients using regression analysis provided by Excel (Microsoft Office Excel 2007).

3 Results

3.1 Gender-specific survival

Table 1 shows the results of the literature research for gender-specific survival. We examined 27 tumors that occur in both sexes, and seven gender-specific malignancies. According to the number of studies, we characterized the gender-specific prognosis of the individual tumors. Furthermore, we recorded the 5-year survival rates.

It is apparent that for more than 50% of the tumor entities, the studies showed a significantly better survival for women. For instance in malignant melanoma, gastric cancer and esophageal cancer, all studies reported a survival advantage of women. Also in lung cancer, all studies except one indicated a better prognosis of women.

But there were also entities in which the majority of studies have noted no difference in survival, e.g. in urinary bladder carcinoma and osteosarcoma.

In a third group of tumors the available data was not conclusive, the number of trials with similar survival times in men and women were equal to those reporting a survival advantage of women. Also in lung cancer, all studies except one indicated a better prognosis of women.

3.2 Tumor ploidy

In this part of the study the total modal chromosome numbers of tumors were analyzed and their frequency distributions represented by a histogram chart (Fig. 1). The data was retrieved from the Mitelman database of chromosomal alterations in cancer. In total, 36,859 karyograms were analyzed. Based on the histograms it became evident which chromosome numbers and thus which ploidy level were most frequently present in the different tumor entities. Representations of colon cancer, chronic myeloid leukemia, osteosarcoma and testicular tumors are shown in Fig. 1. The histograms of all other evaluated tumor entities are available as supplementary data (supplementary figure 1).

All tumor entities, except testicular tumors, showed a clear peak of chromosome numbers in the peridiploid range. In several entities, this was actually the only peak in the distribution. In AML, CLL, salivary gland tumors and thyroid carcinomas the tumors were peridiploid in the overwhelming number of cases. Also CML, ALL, B-cell lymphomas, the fibroblastic/myofibroblastic tumors, liposarcoma, Wilms tumor, as well as multiple myeloma showed a singular diploid peak, but with a somewhat more diffuse distribution and the presence of hyperdiploid and hypodiploid cases. The penis and prostate tumors showed also these peridiploid peaks, but moreover an accumulation of chromosome numbers in the tetraploid range was observed.

Several tumors showed chromosome numbers in the triploid range, as exemplified in the Fig. 1 for colon carcinoma and osteosarcoma. Other tumor types of this group were esophageal carcinoma, gastric carcinoma, gallbladder carcinoma and lung carcinoma. Also pancreatic carcinoma, leiomyosarcoma, as well as ovarian cancer showed a similar distribution which was also detectable in Hodgkin lymphoma, malignant fibrous histiocytoma and cervical cancer.

In addition, there is a group of tumors which had chromosome numbers in the triploid range but without a clear peak in the frequency distribution. This group includes renal tumors, malignant melanoma, bladder carcinoma, hepatocellular carcinoma, GIST, astrocytoma grade III-IV, endometrial tumors, breast cancer and the fibrohistiocytic tumors.

Very unusual is the distribution of chromosome numbers of testicular tumors which are actually never peridiploid but instead showed a peak between the diploid and triploid range (Fig. 1).

We did not analyze the karyograms of 37 tumor types in the Mitelman database for specific karyotypic changes, but it was noted that even within the group of diploid tumors (No. of chromosome=46) there was not a single karyotype without at least one structural or numerical aberration.

3.3 Correlation between ploidy and survival

Using a regression analysis, the extent of chromosomal changes in various tumor entities were compared with the published survival rates. Specifically, the number of cases with hyperdiploid, predominant triploid, predominant hypertriploid and aneuploid chromosome sets were calculated and correlated with the 5-year survival rates. The calculated regression/correlation coefficients showed a significant association between a survival decrease and an increase of aberrant chromosome numbers in carcinomas (Fig. 2). In the analysis of all tumor entities, no statistically significant correlation could be demonstrated.

3.4 Analysis of changes in the sex chromosomes

In Fig. 3, the percentages of mutated sex chromosomes were represented for those tumor entities that occur in both
Table 1 Gender differences in cancer survival based on literature analysis

| Tumor                        | number of studies | $♀ > ♂$ | $♀ \approx ♂$ | $♀ > ♂$ | references | 5-year-survival$^b$ | reference$^b$ |
|------------------------------|-------------------|---------|--------------|---------|------------|-------------------|---------------|
| Better prognosis for women$^a$ | malignant melanoma | 10      | 0            | 0       | [17], [18], [19], [25], [26], [40], [41], [64], [66], [71] | $♀=84\%$, $♀=88\%$ | [66]         |
|                             | gastric carcinoma | 6       | 0            | 0       | [11], [21], [37], [42], [55], [79] | $♀=34.9\%$, $♀=35.7\%$ | [79]         |
|                             | lung carcinoma    | 6       | 5            | 1       | [4], [18], [51], [66], [76], [77] | $♀=15\%$, $♀=18\%$ | [66]         |
|                             | colon carcinoma   | 5       | 5            | 0       | [2], [28], [39], [68], [82] | $♀=49\%$, $♀=51\%$ | [68]         |
|                             | oesophagus carcinoma | 3       | 3            | 0       | [1], [18], [31] | $♀=27.3\%$, $♀=55.4\%$ | [31]         |
|                             | liver carcinoma   | 4       | 4            | 0       | [18], [50], [69], [80] | $♀=16\%$, $♀=22\%$ | [80]         |
|                             | B-cell-lymphoma   | 4       | 4            | 0       | [18], [46], [66], [83] | $♀=62\%$, $♀=66\%$ | [66]         |
|                             | oral cavity carcinoma | 6       | 3            | 2       | [18], [33], [43], [51], [67], [73] | $♀=35\%$, $♀=51\%$ | [43]         |
|                             | Hodgkin disease   | 3       | 3            | 0       | [18], [36], [75] | $♀=86\%$, $♀=90\%$ | [36]         |
|                             | thyroid carcinoma | 3       | 2            | 1       | [56], [66], [74] | $♀=87\%$, $♀=90\%$ | [66]         |
|                             | multiple myeloma  | 2       | 2            | 0       | [18], [38] | $♀=28.5\%$, $♀=33\%$ | [18]         |
|                             | CLL               | 1       | 1            | 0       | [18] | $♀=62.2\%$, $♀=66.4\%$ | [18]         |
|                             | CML               | 1       | 1            | 0       | [18] | $♀=30.5\%$, $♀=36.9\%$ | [18]         |
|                             | AML               | 1       | 1            | 0       | [18] | $♀=12.7\%$, $♀=13.4\%$ | [18]         |
| Prognosis men $\approx$ women$^a$ | bladder carcinoma | 5       | 0            | 4       | [3], [7], [18], [62], [78] | $♀=50\%$ | [78]         |
|                             | salivary gland tumor | 4       | 1            | 3       | [12], [13], [18], [73] | $♀=66.6\%$ | [13]         |
|                             | osteosarcoma      | 3       | 0            | 3       | [14], [27], [61] | $♀=60\%$ | [27]         |
|                             | liposarcoma       | 1       | 0            | 1       | [48] | $♀=83\%$ | [48] |
|                             | leiomyosarcoma    | 1       | 0            | 1       | [49] | $♀=40\%$ | [49]         |
|                             | malignant fibrous histiocytoma | 1       | 0            | 1       | [30] | $♀=51\%$ | [30]         |
| Inconclusive results$^b$    | GIST              | 4       | 2            | 2       | [8], [9], [52], [65] | $♀=69\%$ | [65]         |
|                             | pancreas carcinoma| 4       | 2            | 2       | [18], [35], [66], [81] | $♀=6.4\%$, $♀=7.6\%$ | [18]         |
|                             | kidney tumor      | 4       | 2            | 2       | [18], [24], [32], [66] | $♀=66\%$, $♀=67\%$ | [66]         |
|                             | ALL               | 2       | 1            | 0       | [18], [63] | $♀=24.2\%$, $♀=21.6\%$ | [18]         |
|                             | Wilms tumor       | 2       | 1            | 1       | [27], [45] | $♀=47.3\%$ | [45]         |
|                             | astrocytoma, grade III-IV | 2       | 1            | 1       | [6], [15] | $♀=51\%$ | [6] |
|                             | gallbladder carcinoma | 2       | 1            | 0       | [18], [80] | $♀=6\%$, $♀=11\%$ | [80]         |

$^a$ Evidence for the estimations of gender differences in survival were derived from the number of studies in each tumor type

$^b$ 5-year survival rates as reported by the indicated literature reference. These survival rates were also used for the correlation analysis with the (aneu)ploidy states (see Fig. 2)
women and men. It became evident that the percentage of numerical sex chromosome aberrations is generally higher in men compared to women suggesting a potential association with the gender specific difference in survival. However, when correlating the percentage differences of the mutated sex karyotypes with differences in 5-year survival rates between women and men, no statistically significant relationship could be established.

4 Discussion

4.1 Differences in gender-specific survival

One aim of this study was to create an overview of the gender-specific survival in malignant tumors. One earlier study already presented data for malignant melanoma, soft tissue sarcoma and cancers of the breast, lung, colorectal, esophagus, gastric, pancreatic and soft tissue [47]. Except for pancreas and breast cancer a better survival of women could be demonstrated. Apart from many individual reports which examined gender as a possible prognosis parameter among many others, there are only few comprehensive studies which deal with this issue in multiple tumor entities. Noteworthy in this context is the Eurocare-3-study and for Germany the data of the Robert Koch Institute (RKI).

The Eurocare-3 study examined among other things the gender-specific survival of 42 different malignancies in adults and from 24 malignant tumors in children. Overall, the study included 1.8 million adults and 24,000 children from 20 European countries who were diagnosed with cancer during the period from 1990 to 1994. The patients were observed 5 years after diagnosis until 1999. Various factors were compared with each other, for example the survival between countries, the survival among the different tumors and the survival of women and men. It was recognized that for most tumors, women had a significantly better survival [18]. The results of the Robert Koch Institute represent retrospectively collected data being based on the information from the tumor registries of individual states of Germany. Again, an overall survival advantage of women for most tumor entities could be demonstrated [66]. To our knowledge, this is the first study which primarily focused on this issue and analyzed a large number of tumor types with compilation of data from various reports to provide an estimate of the evidence for gender specific difference in cancer survival.

In the literature the following possible causes for the differences in gender-specific survival are mentioned: differences in tumor biology, defense mechanisms, perception of symptoms and stage at the time of diagnosis. Generally it is stressed that women live healthier, attend
more often screening programs and look for medical advice earlier than men. Thereby tumors may get detected and treated at an earlier stage of the disease.

However, many studies taking into account tumor stage also demonstrated that gender remains a significant and independent prognostic factor. This could be shown, for example, in malignant melanoma [64], colorectal [82] and lung cancer [70]. Thus, it must be postulated that tumor-biological parameters being associated with the sex of the patient are responsible for the survival advantage of women. In addition, the better overall life expectancy of women need to be considered in this context.

4.2 Ploidy and survival

Another objective of this study was to investigate the ploidy level of tumors. This was done by analyzing data from the Mitelman database allowing the assessment of the number of chromosomes in various tumor entities [44]. A similar approach was performed recently by Storchova and Kuffer [72]. These authors, however, placed their emphasis on the analysis of tetraploidy. In our study, particular attention was given to the analysis of triploidy, i.e. tumors exhibiting chromosome numbers in the triploid range, and the question whether this state has any tumor-biological relevance with respect to cancer survival and gender-specific prognosis.

Our analysis indicated that many tumor entities have chromosome numbers with a bimodal distribution with a peak in the triploid range in addition to a diploid one. This was observed particularly in tumors with a high mortality like lung, pancreas, gastric and colon cancer. For lung cancer, we recently showed that a near triploid DNA content is frequently detectable in non-small cell carcinomas [60]. In fact, Ozaki et al. already reported in 1981 that near-triploidy represent the most frequent DNA distribution in lung cancer [54]. Thus, triploidy is not a cell culture artifact. The observation of tripolar mitoses was indicative for a DNA peak in the triploid range [60]. Based on the correlation analysis with survival we could show that the detection and in particular the extent of triploidy is associated with poor survival in carcinomas. Triploidy in this context is obviously correlated with aneuploidy.

In an earlier study by DNA cytometry the bimodal distribution of chromosome numbers was shown for many tumors [5]. The authors already pointed to the influence of the modal DNA value and the ploidy level on prognosis. For the vast majority of tumors, with the exception of squamous cell carcinoma of the cervix uteri, a survival
benefit was observed for the near diploid tumors compared to those in the triploid-tetraploid range [5]. Our study confirmed that the extent of abnormal chromosome numbers in carcinomas is associated with worse prognosis (Fig. 2). This finding again highlights the importance of aneuploidy and chromosomal chaos for cancer survival. In addition it suggests that quantifying the amount of non-diploid cells in carcinomas might be used as a prognostic marker.

The data for non-carcinoma tumor types is heterogeneous. At least there is not such a clear association between the extent of triploidy or aneuploidy with survival. In this context, the analysis of Kaneko and Knudson in neuroblastoma is interesting. It is one of the few reports that dealt with the relevance of ploidy level on tumor behavior. The authors demonstrated that neuroblastoma in lower tumor stages and with improved survival more frequently had a near triploid chromosome set while the near diploid tumors were characterized by advanced stages and poor survival [34].

Tripolar mitosis and triploidy are also detectable in normal cells and are associated with each other as was already shown more than 40 years ago. Tripolar mitoses represent a subtype of multipolar mitoses and occur usually in octaploid or tetraploid cells. During cell division these cells generate triploid and haploid caughter cells (8n: 3n+3n+2n, 4n: 3n+1n). However it is a rare phenomenon that occurred in only 3.6% of the analyzed mitoses [57], [58]. In malignancies, in contrast, cells with a near triploid chromosome content frequently represent the predominant tumor cell clone. And this may be reflected microscopically by the presence of tripolar mitoses. Thus, triploidy probably represents a metastable condition of the chromosome complement.

Our study revealed profound differences in the incidence of triploidy. In some entities it is detectable very often while being almost absent in others. This may have practical importance in the differential diagnosis of malignant undifferentiated tumors of unknown primary site. The detection of a triploid tumor cell clone by DNA measurement or the observation of tripolar mitoses may provide important information in this regard.

Testicular tumors revealed a peculiar histogram of the chromosome number because it did not show the typical peak in the diploid range. This was also observed by DNA cytometry [5]. It can only be speculated whether this distribution has any biological or clinical relevance [23]. In this context we would like to mention that germ cell tumors belong to those entities which frequently show good response to chemotherapy and even spontaneous tumor regression [10]. The histogram suggests that the chromosomes numbers of germ cell tumor may be more frequently in an unstable state between diploidy and triploidy which may be associated with the above mentioned phenomena.

Fig. 3 Analysis of percentage of cases with XX or XY numerical alterations in the different tumor entities (blue colour: percentage of cases in men with XY numerical alterations; red colour: percentage of cases in women with XX numerical alterations)
By analyzing the 36,859 karyograms of the Mitelman database we did not find a single case with normal chromosome number \((n=46)\) without at least one structural or numerical aberration suggesting that chromosomal changes are an inherent feature of cancer formation.

Another observation of interest was the fact that numerical aberrations of the sex chromosomes are more prevalent in tumors from male compared to female patients. The imbalance in relation to the two different gonosomes in men may lead to an increased chromosomal instability. This, in turn, may predispose to the accumulation of additional chromosomal changes as the major cause for tumor progression and cancer related death. Our observation and the above mentioned hypothesis represents a possible biological cause for the phenomenon of worse survival in men. However, it should be noted that the difference in the number of changes of the sex chromosomes in the different tumor entities could not be significantly associated with the difference in the gender-specific survival rates. Apart from alterations of the sex chromosomes, other factors probably play a role within individual tumor types.

In conclusion, a significant survival advantage of women could be shown for most cancer types. Triploidy is a common phenomenon in many but not all tumor entities, its extent is associated with poor prognosis in carcinomas. Tumors of male patients more frequently harbour numerical alterations of the sex chromosomes than tumors of females.

We suggest that the biological mechanisms underlying gender specific differences in survival should be further investigated. Also the role of triploidy in cancer biology and progression merits more attention. Finally, our study again highlights the fundamental importance of chromosomal changes in tumor biology.

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