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Original article

Rapidly and slowly growing lineages in chromosomal instability-type gland-forming gastric carcinomas as revealed by multisampling analysis of DNA copy-number profile

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Short running head: Loss-rich and gain-rich lineages in GC

Abstract

Background: To examine whether gastric carcinoma (GC) with chromosomal instability (CIN-type GC), the largest category in the Cancer Genome Atlas classification, consists of a single genetic lineage, we conducted a multisampling analysis of genomic DNA copy-number profile. *Methods*: We performed array-based comparative genomic hybridization using formalin-fixed, paraffin embedded tissues from 54 gland-forming GCs containing a total of 106 DNA samples from mucosal, extra-mucosal invasive, and lymph node lesions. Microarray data were analyzed by unsupervised hierarchical clustering and penetrance plots. Epstein-Barr virus infection status and mismatch repair (MMR) enzyme silencing/p53/mucin expression were examined with in situ hybridization and immunohistochemistry, respectively. *Results*: The samples examined were divided into gain-rich clusters A and loss-rich cluster B, which were different in tumor locus and patient age. The T1/T2-4 ratio, the frequency of small cancers (diameter $\leq 2-4$ cm), and intestinal mucin expression were higher in cluster B than in cluster A, whereas there were no significant differences in the frequencies of MMR silencing, mutant p53 pattern, and lymph node metastasis between the two clusters.

Conclusions: We demonstrated that the CIN-type GC could be categorized into two genetic lineages which were different in rapidity of local extension but similar in nodal metastasis risk.

Keywords

Stomach, adenocarcinoma, chromosomal instability, copy-number alterations, array-based comparative genomic hybridization, metastasis risk

Introduction

Gastric carcinoma (GC) is the fifth most common malignant neoplasm and the third leading cause of cancer-related deaths in both sexes worldwide [1]. Recently, the Cancer Genome Atlas (TCGA) classification of GC includes four subtypes: Epstein-Barr virus (EBV)-positive GCs, microsatellite-unstable GCs, genomically stable GCs, and GCs with chromosomal instability (CIN) [2]. It remains unclear how the risk of progression from early to advanced stage or metastasis risk is assessable and whether the most common CIN-type GCs are genetically homogeneous.

Early GC is defined as a tumor that is limited to the mucosal and submucosal layers, if present, irrespective of the lymph node status. In Japan, early GCs account for approximately 40%–60% of all GC cases [3-5], most of which are detected by endoscopy. Endoscopic resection (ER) of early GCs, which has become the standard treatment in Japan, is gaining greater acceptance worldwide [6]. However, indications for ER of early GC are limited to those with a very low risk of lymph node metastasis; gland-forming tumors ≤ 2 cm at clinical pT1a stage [7]. When the ESD specimen is pathologically assessed as non-curative, additional treatment follows to prevent recurrence. However, the subsequently removed gastric tissue frequently do not show any evidence of tumor spread or metastasis.

The extent to which early detection and treatment contribute to the reduction of mortality in these patients depends on the lineage continuity between the endoscopically resectable lesions and advanced cancer. In our previous studies, we applied genomic DNA copy-number alteration (CNA) profiling to precursor lesions as well as early and advanced GC specimens, and classified GC samples using unsupervised hierarchical cluster analyses. Using this approach, we have demonstrated that virtually all undifferentiated (diffuse) early GC cases were considered to become advanced [8], whereas around 20% of non-invasive (glandforming) neoplasms were considered to eventually become invasive [9]. In the present study, we utilized this approach to multiple samples from mucosal, extra-mucosal invasive, and metastatic lymph node lesions of individual tumor specimens to confirm the consistency of individual-specific changes and to examine changes associated with tumor progression.

Patients and Methods

Materials

This study used formalin-fixed paraffin embedded tissue specimens from 57 invasive gastric adenocarcinomas including 22 intramucosal, 6 submucosal (including 1 collision cancer, counted as 2 distinct tumors), and 29 advanced cancers (including 2 double cancers, each counted as 2 tumors) (online suppl. Table 1). Of these tumors, 25 lymph node-positive (N+) and 22 node-negative (N0) tumors were surgically resected from 44 patients in the period between 1998 and 2014, and 10 mucosal GCs (cases M27 to M36; online suppl. Table 1) were removed by ESD during the 2012–2014 period. For diagnosis of N0 tumors, 10 or more lymph nodes examined had to be free from metastasis [10]. We performed multisampling, including mucosal, invasive, and if present, lymph node metastatic samples in all 27 advanced GC cases included in this study. The third edition of the Japanese Classification of Gastric Carcinoma and pTNM staging were used to determine histological characteristics and tumor stages, respectively [11].

Immunohistochemistry and EBV in situ hybridization

Immunohistochemical staining of 4- μ m-thick paraffin sections was performed using an automated Ventana Discovery XT system (Tucson, AZ, USA) with heat pretreatment, an

amplification kit (Ventana, 760-080) and a DAB detection kit (Ventana, 760-124). The following two monoclonal antibodies against two mismatch repair (MMR) proteins were used to assess enzyme silencing, which is closely related to microsatellite instability (MSI) [12]: MSH6 (clone 44, Ventana) and PMS2 (clone EPR3947, Cell Marque, Rocklin, CA, USA). A monoclonal antibody to p53 protein (DO-7, 1:100; Dako, Glostrup, Denmark) was used to assess p53 expression pattern. Mucin phenotype was analyzed immunohistochemically using monoclonal antibodies against MUC2 (clone MRQ-18, Cell Marque, Rocklin, CA, USA), MUC5AC (clone MRQ-19, Cell Marque), MUC6 (clone MRQ-20, Cell Marque) and CD10 (clone 56C6, Dako, Glostrup, Denmark). The stains were scored according to the percentage of stained neoplastic cells and categorized into gastric (G), intestinal (I), null (N), G>I, and I>G phenotypes based on the previous study [9]. A polyclonal antibody to SEMA3E protein (1:500; Atlas Antibodies, Bromma, Sweden) to evaluate the expression of SEMA3E gene.

Loss of MMR enzyme expression (LOM) was defined as complete absence of tumor nuclear staining in specimens with retention of MMR enzyme expression (ROM) in nuclei of normal glands and lymphocytes (internal control) [12]. p53 staining pattern was classified as diffuse, regional, sporadic, and null. The former two and the last patterns were considered to reflect mutation, whereas sporadic pattern was considered as a wild-type (WT) pattern. EBV status was determined by EBV-encoded small RNA (EBER) *in situ* hybridization [13] using the INFORM EBER Probe (Ventana, 800-2842: Mannheim, Germany).

Genomic DNA extraction

Genomic DNA was extracted from 5-µm-thick tumor and normal gland (reference) sections using laser microdissection (LMD6000; Leica Microsystems, Wetzlar, Germany). Samples were obtained from an area of \geq 6 mm² in which \geq 90% of the cells were neoplastic. A proteinase K solution (200 µg/ml) was used for digestion of the dissected tumor and reference samples for 70 ± 2 h at 37°C, followed by phenol/chloroform DNA extraction. DNA quality assessment was based on a cut-off A260/A280 ratio of >1.5, a cut-off A260/A230 ratio of >1.0, and concentration of double-stranded DNA.

Whole genome amplification

We used the GenomePlex whole genome amplification kit (WGA2 Kit; Sigma, St. Louis, MO, USA) for DNA amplification according to the manufacturer's protocol [14].

Array-based comparative genomic hybridization

For array-based comparative genomic hybridization (aCGH), 60-mer length oligonucleotide probes were used according to the manufacturer's guidelines [15]. From each tumor specimen, a set of genomic DNA samples, one from nonneoplastic glands, as a reference, tumor and the others from tumor parts, were labeled using Cyanine 5 and Cyanine 3, respectively, prior to competitive microarray hybridization (SurePrint G3 CGH Microarray 8x60K, GPL10152 62,976 probes). Intensity of all hybridized probes were captured and qualified by a DNA microarray scanner (Feature Extraction software 10.7.3.1) followed by calculation of the ratio of tumor and reference fluorescence intensities. Next, chromosomal patterns within the microarray profiles were visualized, detected, and analyzed by the Agilent CGH analytic software using the UCSC Genome Browser according to the latest resource content: hg19 assembly-Design ID 021429 (GRCh Build 37). Definition of genomic copy number gain and loss, and amplification were based on base 2 logarithm of the tumor-toreference (T/R) ratios which were >0.3219, <-0.3219, and >1, respectively. The microarray data were registered in the Gene Expression Omnibus (GEO) data base (Accession number: GSE108507).

Clustering algorithm

Before performing the cluster analysis, average T/R ratio of the probes within each gene was calculated to intensify the signal-to-noise ratio in hybridization analysis. Given that samples from the same tumor might share random generation of gene CNAs and tissue environmental selection, different parts of a tumor might have similar CNA profiles in the presence of a common carcinogenetic process [21, 22]. CNA profiles that have a low noise level could

satisfy this internal standard. Gene size, which is defined as the number of probes in each gene, impacts noise-canceling function of the average of T/R ratio, whereas clustering reproducibility depends on the number of genes. For smaller gene numbers, the reproducibility of clustering analysis becomes lower, whereas gene size is larger. Thus, for determining optimal gene size and number, the clustering analysis was repeatedly performed using the Clustering 3.0 (version 1.52) software and TreeView (version 1.1.6r2) [29, 30]. The unsupervised clustering analysis was based on genomic copy number profile resemblance, and complete linkage and uncentered correlation distance were applied.

Genes showing significantly different CNAs between clusters

Welch's *t* test was used between average $\log_2(T/R)$ values of total A cluster and total B cluster samples for 14,977 protein-coding genes. Bonferroni correction was used to correct for multiple comparisons.

Statistical analysis

We used Microsoft Office Excel 2013 and BellCurve for Excel (Social Survey Research Information Co., Ltd., Tokyo, Japan). Correlations among variables were examined using Fisher's exact test and Welch's *t* test. Statistical significance was determined at a *p* value of less than 0.05.

Results

Unsupervised clustering analysis of the CNA profile

To improve the CNA signal-to-noise ratios, probe T/R ratios within a specified gene were averaged (T/R ratios of 55,142 probes in 30,471 gene regions). To classify the samples solely by the similarity in their CNA profile, unsupervised cluster analysis was performed. To determine the optimal clustering condition, we repeated the clustering nine times using varying gene sizes, from 9,487 genes with ≥ 2 probes to 370 genes with ≥ 10 probes, [9] and correlated the neighboring or splitting sample distribution in the dendrogram with the presence of large chromosomal changes common to all samples for each case, which we defined as stemline changes. For this purpose, we prepared penetrance plots of individual samples (online suppl. Fig. 1). Of these nine conditions, we chose the condition including genes with ≥ 5 probes because the neighboring sample distribution in the dendrogram was best correlated with the presence of stemline changes. In the chosen condition, out of the 27 cases included in the multisampling, 21 cases showed a neighboring pattern and 6 cases showed a splitting pattern in the sample distribution (Fig. 1). Of the 21 neighboring pattern cases, stemline chromosomal changes were detected in 17 cases with ROM and not in 4 cases with LOM (online suppl. Fig. 1). Of the six splitting pattern cases included two double

cancers (case# A13 and A17), three cancers with almost no common chromosomal changes in the samples (case# S5, A15, and A29), and one cancer with stemline changes but splitting between the samples with and without 1p/q+ (case# A7).

Clustering results and their relationship to MMR enzyme expression, p53 expression, EBV infection, and mucin phenotype

Consequently, a total of 106 samples were classified into two major clusters: A and B (Fig. 1). The heat map indicated that there was a copy-number gain area common to both clusters. Clusters A and B were characterized by the presence of gain-rich and loss-rich areas, respectively. LOM samples characteristically showed dark in the heat map. There were no differences in the frequencies of LOM and mutant p53 pattern between the two clusters. The LOM samples were not distributed in the clusters. Additionally, we confirmed that MMR enzyme silencing and mutant p53 pattern were mutually exclusive (Fig. 1). The tumors we examined in this study contain only 1 case with EBV infection. The staining results of mucin phenotype are shown in online suppl. Table 1. We found that the relative frequency of gastric predominant phenotype (G, G≥I)/null type (N) was significantly lower in invasive/metastatic parts than in mucosal parts, whereas intestinal predominant phenotype (I, I>G) was almost constant between these parts (online suppl. Fig. 2). Therefore, we can tentatively regard the intestinal expression as a progression-independent lineage marker. This marker expression

was more common in cluster B than in cluster A.

Chromosomal CNAs

The penetrance plots of the tumors with LOM demonstrated infrequent gains (such as 12q+, which was rare in the ROM samples) and scarce losses, whereas gains and losses were conspicuous in the ROM tumors (Fig. 2a, b), which were divided into two patterns by clustering (Fig. 2c, d). The changes common to both clusters were 8q+, 13q+, 20q+, and 5q-. Clusters A and B were characterized by gain-rich (7p/q+, Xp/q+, etc.) and loss-rich (4p/q-, etc.), respectively (Fig. 2c, d). Statistical analyses are shown in Table 1a. The mucosal, invasive, and metastatic ROM samples were compared in clusters A and B (Fig. 3). The changes common to clusters A and B (8q+, 13q+, 20q+, and 5q-) were already present in the mucosal samples, whereas cluster-specific changes (such as 4p/q- and 7p/q+) accumulated during the progression to invasion and metastasis. These were confirmed by statistical analysis of the frequency of each chromosomal CNA, which were determined by counting using the penetrance plot of each sample (online suppl. Fig. 1) and compared between the mucosal and extra-mucosal (invasive and metastatic) samples (Table 1b). In contrast, such differences were not detected between the invasive and metastatic samples. Table 1c shows metastasis-related chromosomal changes. The chromosomal changes common to clusters A and B were not or weakly related to metastasis, whereas the cluster specific changes, such as

7p/q+ in cluster A and 3p-, 4p-, 5q-, and 8p- in cluster B were significantly related to lymph node metastasis. The chromosomal changes detected are summarized as an evolutionary tree in Fig. 4.

Clinicopathological and molecular characteristics of the clusters

Clinicopathological characteristics and molecular changes were compared between the tumors of clusters A and B (Table 2). The tumor locus and patient age were different between clusters A and B. In this tumor-based analysis, representative phenotype in each submucosal or advanced cancers was defined to that of the invasive sample, and we could also confirm that the intestinal expression was more frequent in cluster B than in cluster A.

The T1/T2–4 ratio and the frequency of small cancers (diameter $\leq 2-4$ cm) were higher in cluster B than in cluster A, suggesting that the GCs in cluster A is more rapidly growing than in cluster B. Nine of the 10 ESD specimens examined were classified into cluster B. There were however no significant differences in the frequencies of lymph node metastasis.

Genes showing significantly different CNAs between the clusters

We identified 32 genes that significantly contributed to the difference in clusters A and B. Of these genes, 12 were related to growth, as shown in online suppl. Fig. 2. When the gene function and the direction of CNAs were consistent, we regard the CNAs as putative driver

and when inconsistent, we regard the CNAs as putative passenger. There were five putative driver tumor suppressor genes (*CACNA2D3*, *PTPRG*, and *LRIG1* at chromosome 3p, *SLIT2* at chromosome 4p, and *FSTL5* at chromosome 4q) and one possible driver protooncogene (semaphorin 3E [*SEMA3E*]) at 7q. However, immunohistochemically, there was no difference in expression level of SEMA3E protein between 10 tumors that showed greatest copy-number gains and those with greatest copy-number losses.

To compare the growth activity between cluster A and cluster B tumors, we examined the frequency of amplification of growth-related genes, which is a characteristic of CIN-type GC. Of the 55 receptor tyrosine kinases (RTKs) examined, copy-number gains of 12 RTKs, including EGFR, ERBB3, FGFR4 and EPHB3 and amplification of LMTK2, EPHB3, EPHB4 showed significantly different frequency between clusters A and B, and all of them were more frequently detected in cluster A than in cluster B (online suppl. Table 2).

Discussion

Using the multisampling method, we could demonstrate not only the differences among samples but also the reproducibility of changes common to all samples in individual GC patients. To assess the reproducibility of this approach as well as to classify samples, we applied unsupervised, hierarchical clustering analysis to the multisampling aCGH data of 57 early and advanced gland-forming GC specimens. We selected the clustering condition that showed the highest concordance between the neighboring pattern in the clustering dendrogram and the presence of stemline changes. Even in the absence of stemline changes among the samples from individual tumors such as the LOM tumors, the mucosal and invasive/metastatic samples of individual tumors often showed a neighboring position in the dendrogram, likely reflecting a similarity among gene-level CNAs. CNA profiles are considered as individual tumor-specific and progression-independent lineage markers.

During development of individual tumors, random changes as well as essential genetic and epigenetic changes accumulate in a time-dependent manner based on genetic instability, and these changes undergo natural selection by the tissue environment. In this evolutionary process, genomic alteration profiles might become unique to each individual tumor, which might converge in several types by natural selection [16]. We attempted to reveal such converged genotypes using unsupervised hierarchical clustering.

The tumors included in the present analyses were mostly CIN-type and partly MSI-type of TCGA classification, whereas EBV-related GCs were infrequent. A cancer with LOM, which largely overlaps with the MSI phenotype, is characterized by a low frequency of chromosomal changes and the presence of the unique 12q+ (Fig. 2a, b), and may have branched off from the main trunk of gland-forming GCs at earlier stages of GC development (Fig. 4). Accordingly, it was reported that GCs of MSI phenotype showed the frequency of chromosomal CNAs lower than in GCs without MSI and were associated with high DNA methylation status even at intramucosal stages [17].

We classified ROM GC samples (GCs excluding LOM) into two clusters, A (ROM) and B (ROM) (Fig. 2c, d). Based on the resemblance of gene-level CNA profiles, our clustering approach have successfully disclosed two distinct genetic lineages (gain-rich and loss-rich lineages) that have not been clearly recognized thus far. In the present study, we analyzed sequential accumulation of chromosomal CNAs during progression from the mucosal to the invasive/metastatic growth in each lineage (Fig. 3). Both lineages might have derived from the common trunk with 8q+, 13q+, 20p/q+, and 5q–,which were commonly detected in both clusters A and B. These findings partly confirm the report of Uchida *et al.*, who reported 8p+ and 5q– as early changes of GC [18]. Then, cluster-specific later changes, such as 7pq+ in cluster A and 4pq– in cluster B (Fig. 4).

Next, we classified all samples into two clusters, A and B. There appeared little relationship between this cluster classification and TCGA GC subtypes (Table 2). The tumor locus and patient age were different between clusters A and B. The age difference might be explained by the difference in the frequency of early GCs, whereas the locus difference suggests that this clustering might reflect distinct genetic lineages. This was also supported by mucin phenotyping; progression-independent intestinal expression was significantly different between clusters A and B. Comparison of other clinicopathological factors between clusters A and B indicated that the tumors in cluster A was more rapidly growing (larger, deeper tumors) than those in cluster B. ESD specimens were mostly included in the latter. These finding implied that early detection and treatment were more difficult in cluster A than in cluster B tumors (Fig. 5).

A rapid growth in the gain-rich lineage (cluster A) might be partly due to the copy-number gains/amplifications of the RTKs mentioned above and growth-related genes in chromosome 7, including *RalA* [19], *EGFR* [20], *MAFK* [21], *GLI3* [22], *CUL1* [23], and *SEMA3E* [24, 25] (online suppl. Table 2). Among these, only *SEMA3E* exhibited significant copy-number differences after Bonferroni correction between clusters A and B. However, its expression is reportedly enhanced [24] or suppressed [25] in GCs. Our immunohistochemical studies for *SEMA3E* failed to demonstrate significant association between protein expression level and gene copy number, and we cannot regard this gene as functional driver or suppressor. Rather, detection of immunoreactivity for amplified growth-related genes may be more promising as cluster A markers. In the loss-rich lineage (cluster B), the above-mentioned tumor suppressor genes on 3p and 4p/q showed significant differences between clusters A and B. In particular,

silencing of *CACNA2D3* [26] and *PTPRG* [27] were reported to play a role in gastric tumorigenesis.

There was no significant differences in the lymph node metastasis risk between the rapidly and slowly growing lineages, whereas there were lineage-specific, invasion/metastasis-related chromosomal changes. These findings suggest that metastasis risk is determined not at an earlier but at a later stage during tumor development as shown in Fig. 4.

These conclusions have several clinical implications for early detection and treatment strategies. This study was conducted with a null hypothesis that there are indolent cancers treated as cancers, which are associated with very low mortality risk. We previously demonstrated that high-grade and low-grade gastric adenomas, of which the former is treated as carcinoma in Japan, were such tumors. Only around 20% of these tumors were inferred to progress to overt GC [9] (Fig. 5). The present study showed that there was no inherently indolent CIN-type early GC cluster and that the two lineages demonstrated in the present study had equal potential for metastasis. Our findings also demonstrated that GCs detected at an early stage were biased to a slowly growing loss-rich lineage. The prevalent use of early detection by endoscopy and subsequent treatment contribute to the reductions in the incidence of advanced GC and age-adjusted mortality rates; however, these reduction sizes were smaller than expected compared with the reduction in the GC morbidity rate [28]. One major cause might be the increase in the percentage of older patients. Another factor might be the failure of early detection of GCs with a rapidly growing gain-rich lineage, which remains a challenge that should be addressed in future studies.

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Statement of Ethics

All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1964 and later versions. The Institution Review Board on Medical Ethics at Shiga University of Medical Science granted permission for conducting this study (Permission number: 30-021). A substitute of written informed consent was obtained from all patients included in the study.

Conflict of Interests

The authors declare that they have no conflict of interest.

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Figure legends

Fig. 1: Unsupervised clustering analysis of 106 samples using the genes/markers of \geq 5-probe size. Genomic copy-number gains and losses of genes/markers are indicated by red and green squares, respectively in the clustering heat map. The clustering results are related to tumor size (gray squares: \leq 2 cm), presence of lymph node metastasis (marked with green background of sample names), Mutant pattern (M, purple) of p53 immunohistochemistry (WT; Wild type), loss of mismatch repair enzyme expression (LOM, pink), and EBV infection status, and mucin phenotype (I/G/N: intestinal/gastric/null). Intestinal predominant (I, I > G) and gastric predominant (G, G \geq I) phenotypes are marked with yellow and brown background, respectively.

Fig. 2: Frequency of copy-number alterations at the chromosome level (penetrance plots) compared between the tumors with and those without mismatch repair enzyme silencing and between clusters A and B. Statistically significant differences are marked with blue frames. a,
b. LOM: loss of mismatch repair enzyme expression; ROM: retention of mismatch repair enzyme expression. The sample numbers of LOM and ROM are 14 and 92, respectively. c, d. Clusters A and B consist of 41 and 51 samples, respectively.

Fig. 3: Penetrance plots of ROM tumors in clusters A and B. Cluster-specific, progressionrelated significant changes are marked with blue frames. **a**, **d**. Mucosal samples; **b**, **e**. extramucosal invasive samples; **c**, **f**. lymph node metastasis samples. The sample numbers of **a** to **f** are 18, 15, 8, 31, 12, and 8, respectively.

Fig. 4: Evolutionary tree of gland-forming gastric carcinoma reconstructed from the data of Table 1, and Figs 2 and 3. LOM/ROM: loss/retention of mismatch repair enzyme expression. Chromosomal gains and losses are shown as red and green characters. Bold/normal letters indicate frequent/infrequent significant changes. Chromosomal parts in parentheses indicate insignificant but characteristic changes.

Fig. 5: Progression model of gland-forming gastric neoplasm from intramucosal neoplasm to invasive carcinoma.

Online suppl. Table 1: Clinicopathological features of individual tumors (age, gender, locus, size of mucosal lesion, pT, pN, and excision type) and immunohistochemical and in situ hybridization results (p53, MMR enzymes, EBV, mucin markers) of individual samples.

Online suppl. Table 2: Frequencies of copy-number gains and amplifications of receptor tyrosine kinases and representative growth-related genes on chromosome 7 in clusters A and B. (xlsx15KB)

Online suppl. Fig. 1: Penetrance plots of individual samples. The sample orders in clusters A and B and LOM corresponds to the sample order in Fig. 1. LOM: loss of mismatch repair enzyme expression; ROM: retention of mismatch repair enzyme expression. (pdf282KB)

Online suppl. Fig. 2: Comparison of mucin phenotype composition between mucosal (M) and invasive/metastatic (I + LN) samples in the cluster A and the cluster B tumors. Based on the online suppl. Table 1, mucin phenotype was classified into gastric predominant phenotype (G, G≥I), intestinal dominant phenotype (I, I>G), and unclassifiable phenotype (N). **a.** mucin phenotype composition of M and I + LN parts in clusters A and B. **b, c**. Statistical analyses of the differences in frequency of gastric predominant expression (b) or intestinal predominant expression (c) between M and I + LN parts and between clusters A and B. (pdf389KB)

Online suppl. Fig. 3: Outline of 32 genes that showed significantly different copy-numbers

between clusters A and B. The mean copy number alterations (CNAs) are expresses as Tumor/Reference (T/R) signal intensity ratio. In the gene function column, tumor suppressor genes and proto-oncogenes are marked with green and pink background, respectively. In the mean T/R ratio columns, copy-number gains and losses are marked with pink and green background, respectively. Concordant pairs of CNA and gene function are marked with black frame. (xlsx15KB)

		а			b		С			
	Number o	f samples	p value	Number o	of samples	p value	Number	p value		
	Cluster A $(n = 41)$	Cluster B $(n = 51)$		$M \\ (n = 53)$	I+LN (<i>n</i> = 39)	M vs I+LN	N1-3 (<i>n</i> = 51)	N0 (n = 41)	N1-3 vs N0	
3p—	7	14	0.3192	6	15	0.0028	17	4	0.0114	
3q+	10	6	0.1660	6	10	0.0967	11	5	0.2792	
4p—	4	12	0.1020	5	11	0.0260	14	2	0.0097	
4q-	2	14	0.0051	8	8	0.5820	13	3	0.0275	
5p+	12	8	0.1339	8	12	0.0805	14	6	0.2035	
5q-	8	16	0.2374	11	13	0.2306	19	5	0.0083	
7p+	23	7	< 0.0001	9	21	0.0003	24	6	0.0015	
7q+	12	2	0.001	3	11	0.0064	13	1	0.0025	
8p-	6	16	0.0853	7	15	0.0067	19	3	0.0011	
8q+	19	23	1.0000	23	19	0.6746	25	17	0.5309	
9p—	7	17	0.0969	9	15	0.0300	18	6	0.0320	
10p+	12	7	0.0763	6	13	0.0175	16	3	0.0047	
13q+	15	18	1.0000	15	18	0.0846	23	10	0.0501	
14q-	0	6	0.0316	2	4	0.3955	5	1	0.2202	
15q+	11	6	0.1033	6	11	0.0565	12	5	0.1877	
16p+	11	3	0.0077	6	8	0.2525	11	3	0.0806	
17p-	2	10	0.0592	6	6	0.7554	9	3	0.2144	
18q-	7	10	0.7938	6	11	0.0565	12	5	0.1877	
19p-	2	12	0.0181	5	9	0.0851	10	4	0.2486	
19q—	1	4	0.3764	1	4	0.1587	2	3	0.6528	
20q+	27	29	0.3996	28	28	0.0847	39	17	0.0011	
22q-	0	11	0.0009	4	7	0.1934	8	3	0.3343	
Xp+	18	5	0.0002	8	15	0.0147	19	4	0.0032	
Xq-	1	5	0.2202	1	5	0.0796	6	0	0.0316	
Xq+	19	8	0.0024	10	17	0.0121	20	7	0.0231	

Table 1: Differences in frequencies of chromosomal copy-number changes between clusters A and B (a), between mucosal and invasive/metastatic samples (b), and between N0 and N1-3 (c) in ROM tumors.

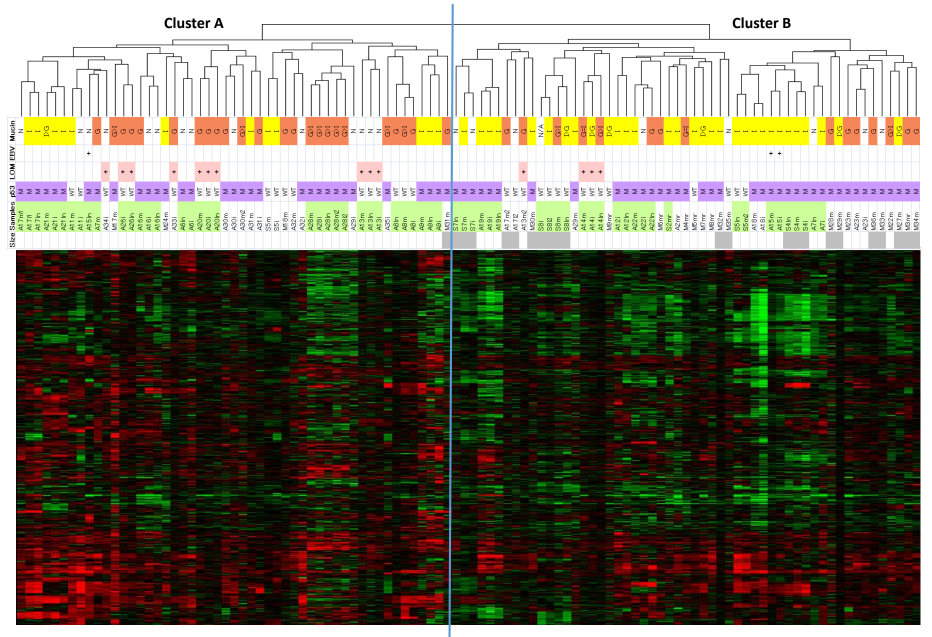
ROM: retention of mismatch repair enzyme expression. *p values* <0.05 are shown in bold.

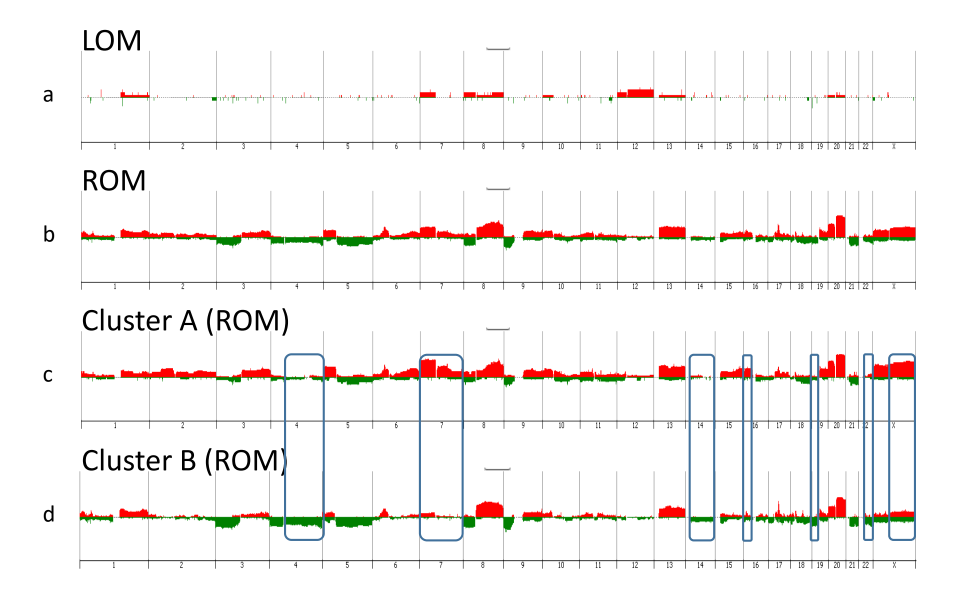
57 tumors	Cluster A	Cluster B	Total	p value
Gender*		Oldblei D	10041	pvalue
- Male	12	22	34	0.5451
- Female	8	9	17	0.0401
Age* (mean \pm SD)	-	68.55 ± 7.24	11	0.006
Tumor locus	10100 - 0100	00100 - 112 1		01000
- Upper (U)	3	9	12	0.0038
- Middle (M)	$\ddot{7}$	19	26	(U+M vs L)
- Lower (L)	13	6	19^{-3}	
Tumor size		-		
≤ 2 cm	1	14	15	0.0019
>2cm	22	20	42	0.0019
≤ 3 cm	5	19	24	0.0143
>3cm	18	15	33	0.0145
≤ 4 cm	9	26	35	0.0062
>4cm	14	8	22	0.0002
$\leq 5 \mathrm{cm}$	14	27	41	0.1454
>5cm	9	7	16	0.1404
Local extension				
T1	5	23	28	0.0011
T2-4	18	11	29	
Nodal lymph node status	10	22	~	
N0	12	23	35	0.2769
N1-3	11	11	22	
Mismatch repair enzyme e LOM	-	2	-	0 1057
ROM	518	$\frac{2}{32}$	7 50	0.1057
EBV infection	10	02	90	
+	0	1	1	1.0000
_	23	33	56	1.0000
p53 expression		00	00	
Diffuse/	10		20	0 1 40 -
regional/null	13	26	39	0.1497
	10	Q	10	
Sporadic	10	8	18	
Phenotype				
Intestinal predominant	5	17	23	0.0264
Non-intestinal	17	15	32	

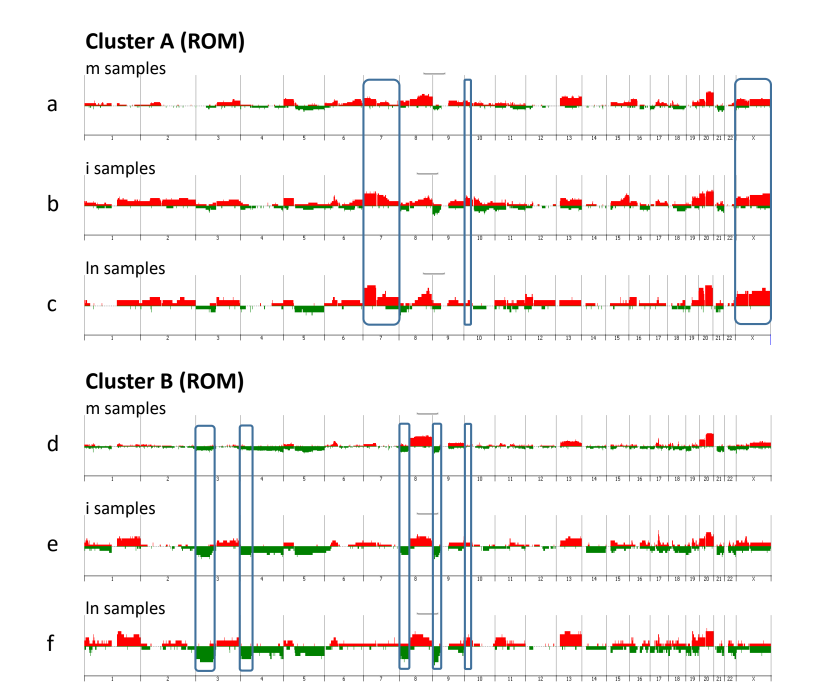
Table 2: Differences in clinicopathological factors between clusters A and B.

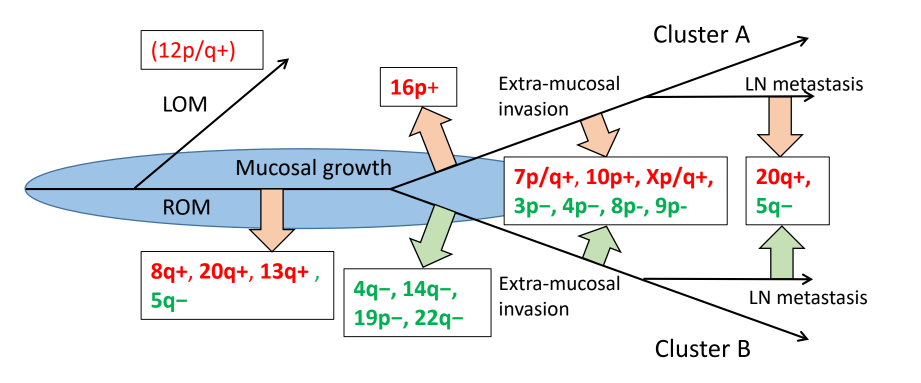
*Tumor number is 51 after excluding 2 double cancers and 1 collision cancer (6 tumors). LOM/ROM: loss/retention of mismatch repair enzyme expression. *p values* <0.05 are shown in bold.

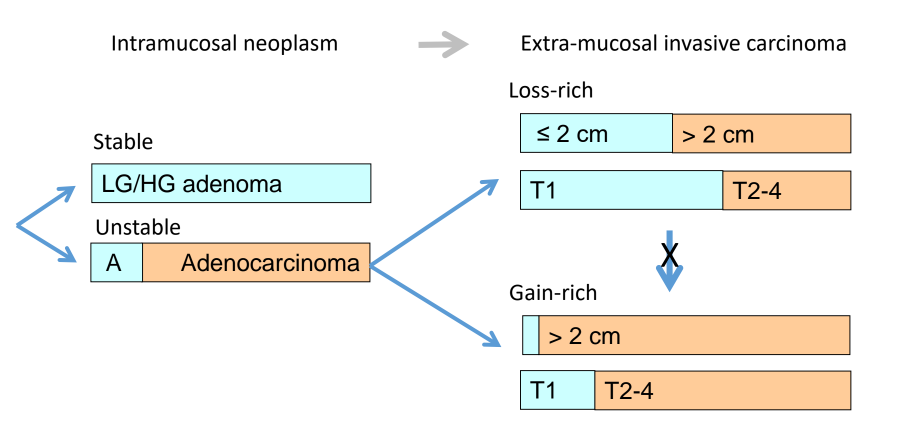
Fig. 1











Online suppl. Table 1: Clinicopathological features of individual tumors (age, gender, locus, size of mucosal lesion, pT, pN, and excision type). and immunohistochemical and in situ hybridization results (p53, MMR enzymes, EBV, mucin markers) of individual samples.

	itu hybridizat I. Samples		Gender	Locus*	Size	<u>- ВV,</u> Т	N	Excision	p53	MSH6	PMS2	LOM†	EBV	MUC5AC	MUC6	MUC2	CD10	Phenotype‡
#1	M3mr	67	F	М	13	1a		Surgical	Null	+	+			2+	—	-	—	G
#2 #3	M4mr M5mr	67 74	M M	M M	30 60	1a 1a	0 0	Surgical Surgical	Diffuse Sporadic	+ +	+ +			1+	_	1+ 1+	— 1+	G=I
#4	M6mr	65	M	M	25	1a	0	Surgical	Diffuse	+	+			1+	_	_	_	G
#5	M7mr	67	F	Μ	25	1a	0	Surgical	Diffuse	+	+			—	1+	1+	1+	I>G
#6 #7	M8mr M9mr	70 51	F M	L	35 40		0	Surgical	Null	+	+			— 1.	_		1+	l I>G
#7 #8	M9mr M17m	51 59	M	L	40 25	1a 1a	0 0	Surgical Surgical	Sporadic Diffuse	+ +	++			1+ 2+	2+	2+ 1+	- 2+	1>G G>l
#9	M18m	80	F	L	25	1a	0	Surgical	Sporadic	+	+			3+	1+	_	_	G
#10	M22m	76	Μ	Μ	22	1a	0	Surgical	Null	+	+			1+	1+	1+	—	G>I
#11 #12	M23m M24m	65 68	M M	M M	35 25	1a 1a	0 0	Surgical Surgical	Null Diffuse	+ +	++			3+	2+	— 1+	— 3+	G
#12	M24m M27m	75	M	M	10	1a	0	ESD**	Diffuse	+	+			_	1+	1+	1+	I>G
#14	M28m	58	Μ	Μ	7	1a	0	ESD	Diffuse	+	+			1+	1+	—	—	G
#15	M29m	62	F	M	7	1a	0	ESD	Diffuse	+	+			1+	—	—	2+	I>G
#16 #17	M31m M30m	81 71	F M	L U	9 10	1b 1a	0 0	ESD ESD	Diffuse Null	+ +	+			2+	_	_	— 1+	G
#17 #18	M32m	61	F	U	9	1a	0	ESD	Null	+	+			_	_	1+	2+	I
#19	M33m	60	Μ	Μ	8	1a	0	ESD	Diffuse	+	+			—	—	—	—	Ν
#20	M34m	61	M	M	7	1a	0	ESD	Regional	+	+			1+	2+	—	—	G
#21 #22	M35m M36m	70 67	M M	U M	5 9	1a 1a	0 0	ESD ESD	Sporadic Diffuse	+ +	++			_ 2+	— 1+	_	_	N G
#23	S2mr	81	F	L	35	1b		Surgical	Diffuse	+	+			_	_	_	1+	
#24	S4m	63	Μ	L	20	1b	1	Surgical	Diffuse	+	+			_	_	—	1+	l
#24 #24	S4i S4ln								Diffuse Diffuse	+ +	++			_	_	_	2+ 2+	I
#24	S5m	82	М	U	35	1b	0	Surgical	Sporadic	+	+			_	_	_	3+	l I
#25	S5i							0	Sporadic	+	+			—	—	—	3+	I
#26	S5m2								Sporadic	+	+			—	—	—	3+	I
#26 #27	S5ln S7m	75	F	М	20	1b	1	Surgical	Sporadic Diffuse	+ +	++			_	_	+	3+ —	l I
#27 #27	57m S7i	23	ı	171	20	тŊ	т	Surgical	Diffuse	+	+			_	_		_	I N
#27	S7ln								Diffuse	+	+			—	—	—	_	Ν
#28 #28	S8m	64	F	Μ	15	1b	1	Surgical	Sporadic Sporadic	+	+			2+ N/A¶	1+ N/A	1+ N/A	— NI / A	G>I
#28 #28	58i 58i2								Sporadic Sporadic	+ +	++			N/A¶ —	N/A 	N/A 	N/A 3+	N/A I
#28	S8In								Sporadic	+	+			_	2+	1+	3+	l>G
#29	A2mr	71	М	Μ	100	3	0	Surgical	Diffuse	+	+			—	—	—	1+	I
#30 #30	A6i A6ln	81	F	Μ	35	3	2	Surgical	Diffuse Diffuse	+ +	+			_	_	_	_	N N
#30 #31	Abin A7m	79	М	U	70	4a	3	Surgical	Diffuse	++	++			 1+	2+	_	_	G
#31	A7i			-			-	0.00	Diffuse	+	+			_	_	—	1+	Ī
#31	A7In		_		. –	-			Diffuse	+	+			_	_	_	—	N
#32 #32	A8m A8i	69	F	L	45	3	3	Surgical	Sporadic Sporadic	+ +	+			2+ 2+	_	1+ —	_	G>I G
#32 #32	Aði Aðin								Sporadic	++	+			2+ 2+	_	_	_	G
#33	A9m	67	F	L	50	4a	2	Surgical	Diffuse	+	+			—	—	3+	2+	I
#33	A9i								Diffuse	+	+			1+	_	2+	1+	I
#33 #34	A9ln A11m	70	F	М	25	2	1	Surgical	Diffuse Sporadic	+ +	++			_	_	2+	— 3+	I
#34	A11i	70	•		20	-	-	Surgicul	Sporadic	+	+			_	_	_	_	N
#35	A12i	65	F	L	55	3	1	Surgical	Diffuse	+	+			—	—	—	3+	I
#35 #36	A12ln A13m	75	F	М	43	2	1	Surgical	Diffuse	+	+	LOM		_	_	_	3+	I
#36 #36	A13ii	75	Г	IVI	45	Z	T	Surgical	Sporadic Sporadic	+ +	_	LOM		_	_	_	_	N N
#36	A13In								Sporadic	+	_	LOM		_	_	—	_	N
#37	A13m2	75	F	U	45	2	0	Surgical	Sporadic	+	—	LOM		1+	_	_	—	G
#38 #29	A14m A14i	77	Μ	L	70	4a	3	Surgical	Sporadic Sporadic	+	_	lom lom		_	3+ 1+	3+ 2+	_	G=I I>G
#38 #38	A14I A14In								Sporadic Sporadic	+ +	_	LOM		3+	1+ 1+	2+ 2+	_	1>G G>l
#39	A15m	71	М	U	95	4a	3	Surgical	Null	+	+		+	_	_	2+	1+	I
#39	A15i								Null	+	+		+	_	_	1+	—	1 N
#39 #40	A15ln A16m	80	М	1	80	3	3	Surgical	Null Diffuse	+ +	++		+	— 1+	— 1+	_	_	N G
#40 #40	A16i	80		L	80	5	5	Surgical	Diffuse	+	+					_	_	N
#40	A16ln								Diffuse	+	+			—	—	—	—	Ν
#41	A17m1	82	Μ	L	40	3	1	Surgical	Diffuse	+	+			—	—	—	-	N
#41 #41	A17i1 A17ln								Diffuse Diffuse	+ +	+			_	_	_	1+ 3+	1
#42	A17m2	82	М	М	15	2	0	Surgical	Sporadic	+	+			1+	1+	_	_	G
#42	A17i2							_	Sporadic	+	+			—	—	—		N
#43 #43	A18m	72	Μ	U	40	3	0	Surgical	Diffuse Diffuse	+	+			—	_	_	3+ 3+	1
#43 #44	A18i A19m	65	М	U	40	3	3	Surgical	Diffuse Diffuse	+ +	++			_	_	+	3+ —	I I
#44	A19i			-	-	-	-		Diffuse	+	+			—	_	+	_	I
#44	A19In			-		~		C	Diffuse	+	+			_	-	1+	1+	
#45 #45	A20m A20i	86	Μ	L	100	3	1	Surgical	Sporadic Sporadic	+ +	_	lom lom		3+ 3+	1+ 1+	_	_	G G
#45 #45	A201 A20In								Sporadic	+	_	LOIVI		3+ 3+	1+ 3+	_	_	G
#46	A21m	79	М	Μ	45	4a	2	Surgical	Regional	+	+			_	1+	—	3+	I>G
#46	A21i								Regional	+	+			—	—	—	3+ 2+	l
#46 #47	A21ln A22m	85	М	М	60	3	3a	Surgical	Regional Diffuse	+ +	+ +			_	_	_ 1+	3+ —	l I
#47 #47	A22ii				20	5	- 4	Broan	Diffuse	+	+			_	_	_	_	N
#47	A22In								Diffuse	+	+			1+	_	—	_	G
#48 #48	A23m	70	Μ	Μ	30	2	0	Surgical	Diffuse Diffuse	+	+			1+ 	_	_		G
#48 #49	A23i A26i	86	F	L	85	3	2	Surgical	Diffuse Sporadic	+ +	+	LOM		2+	_	_	_	N G
#49	A26In			-		-	-	0.001	Sporadic	+	_	LOM		2+	1+	_	_	G
#50	A28m	85	Μ	Μ	95	3	>1	Surgical	Null	+	+			2+	1+	1+	_	G>I
#50 #50	A28i A28ln								Null Null	+ +	+ +			3+ 3+	1+ —	2+ 1+	_	G>I G>I
#50 #50	A28m2								Null	++	+			3+ 2+	1+	1+ 1+	_	G>I
#50	A28i2								Null	+	+			2+	2+	2+	_	G>I
#51	A29m	86	Μ	Μ	31	3	0	Surgical	Diffuse	+	+			—	-	_	2+	1
#51 #52	A29i A30m	78	F	I	48	3	0	Surgical	Diffuse Diffuse	+ +	+ +			— 1+	_	_	_	N G
#52 #52	A30i	70		Ľ	υ	J	5	Surgical	Diffuse	+	+				_	_	_	N
#52	A30m2								Diffuse	+	+			_	2+	1+	_	G>I
#53 #52	A31m	80	Μ	U	56	3	0	Surgical	Regional Regional	+	+			— 2+	_	_	3+	l G
#53 #54	A31i A32m	71	М	L	60	4b	0	Surgical	Regional Sporadic	+ +	++			2+ 3+	_	_	_	G G
#54	A32i	. =		-	_ -		-	0.001	Regional	+	+			_	_	_	_	N
#55	A33i	72	М	L	120	3	0	Surgical	Sporadic	+	_	LOM		1+	2+	_	_	G
#56 #57	A34i A35i	57 72	M M	L U	60 60	3 2	0 0	Surgical Surgical	Sporadic Diffuse	+ +	 +	LOM		 3+	— 3+	_	— 1+	N G>l
								-				oction t						

*Locus: "U"/"M"/"L", upper/middle/lower one third of the stomach. **ESD: endoscopic submucosal dissection. +LOM: Loss of mismatch repair enzyme expression. + Phenotype: "G"/"I"/"N", gastric/intestinal/null. ¶N/A: not assessable.

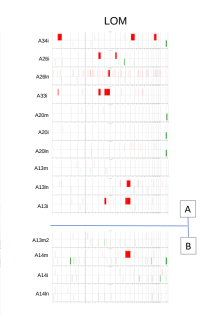
		py-number gain			nplification		
Name of Gene	Cluster A	Cluster B	p value	Cluster A	Cluster B	p value	
	(n = 51)	(<i>n = 5</i> 5)	p vulue	(n = 51)	(<i>n = 5</i> 5)	p vulu	
AATK	8	13	0.3388	1	0	0.481	
ALK	6	4	0.516	0	0	1	
AXL	15	3	0.0014	1	0	0.481	
CSF1R	10	12	0.8148	0	1	1	
DDR1	20	19	0.6887	3	3	1	
DDR2	16	19	0.8368	2	0	0.229	
FGFR1	13	12	0.8193	2	1	0.607	
FGFR2	5	12	0.1155	0	1	1	
FGFR3	21	17	0.314	8	4	0.224	
FGFR4	17	8	0.0381	3	1	0.349	
FLT1(VEGFR1)	25	19	0.168	5	1	0.103	
FLT3	16	23	0.3158	3	0	0.107	
FLT4 (VEGFR3)	19	14	0.2131	1	0	0.481	
IGF1R	16	6	0.0154	2	0	0.229	
INSR	3	2	0.6699	0	0	1	
INSRR	N/A	N/A	N/A	N/A	N/A	N/A	
KDR (VEGFR2)	4	2	0.4248	1	0	0,.81	
кіт	6	3	0.3075	1	0	0.481	
LMTK2	29	15	0.003	7	0	0.004	
LMTK3	N/A	N/A	N/A	N/A	N/A	N/A	
LTK	N/A	N/A	N/A	N/A	N/A	N/A	
MERTK	14	12	0.6519	0	1	1	
MET	18	11	0.0861	0	1	1	
MST1R	17	7	0.0191	1	0	0.481	
MUSK	13	9	0.3382	0	0	1	
NTRK1	13	17	0.6667	1	1	1	
NTRK2	9	7	0.5901	0	0	1	
MUSK MUSK NTRK1 NTRK2 NTRK3	14	4	0.0086	0	0	1	
	4	3	0.7086	0	0	1	
PDGFRB	7	8	1	0	0	1	
PTK7	22	21	0.6932	13	10	0.48	
RET	19	19	0.8405	2	7	0.163	
ROR1	5	6	1	0	0	1	
PDGFRA PDGFRB DGFRB PTK7 RET ROR1 ROR2	7	3	0.1905	0	0	1	
ROS1	10	2	0.0129	0	0	1	
RYK	9	3	0.0661	0	0	1	
STYK1	29	20	0.0508	8	3	0.114	
TEK	3	3	1	0	0	1	
TIE1	13	22	0.1484	5	2	0.257	
TYRO3	17	15	0.5313	1	1	1	
EPHA1	6	7	1	0	0	1	
EPHA2	16	, 14	0.5248	0	0	1	
EPHA3	10	4	0.0248 0.0086	0	0	1	
EPHA4	9	5	0.2544	0	0	1	
EPHA5	3	8	0.2344	0	0	1	
EPHA5 EPHA6		ہ 5		0	0		
EPHA6 EPHA7	6 3	2	0.755 0.6699	0	0	1 1	
EPHA7 EPHA8	3 6	12	0.8699	2	0	0.229	
			0.2018	11			
EPHA10	26	29			14	0.655	
	9	5	0.2544	0	0	1	
EPHB2	21	24	0.8457	2	1	0.607	
EPHB3	24	15	0.0444	8	2	0.046	
EPHB4	28	20	0.0786	14	5	0.021	
EPHB6	32	27	0.1755	14	7	0.086	
EGFR	20	7	0.0033	6	1	0.053	
ERBB2	28	32	0.8449	7	13	0.221	
ERBB3	17	8	0,.0381	7	3	0.190	
ERBB4	6	1	0.0537	0	0	1	

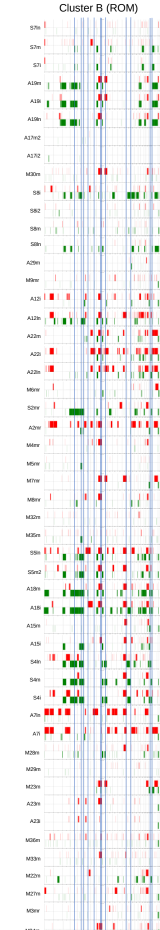
Online suppl. Table 2: Frequencies of copy-number gains and amplifications of receptor tyrosine kinases and representative growth-related genes on chromosome 7 in clusters A and B.

RALA	18	9	0.0283	2	0	0.2291
PTPN12	16	9	0.1079	4	0	0.0503
	34	25	0.0328	23	15	0.0693
	17	15	0.5313	5	3	0.4772
s a CAV1	11	9	0.6206	1	0	0.4811
ထိ ဆို CAV1	12	8	0.3213	1	0	0.4811
GLI3 HOXA1 HOXA1 GLI3 HOXA1 GLI3 GLI	25	11	0.0021	1	0	0.4811
HOXA1 4 4 5 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	29	32	1	4	3	0.7086
E S RABL5	21	13	0.063	4	3	0.7086
စ္	22	21	0.6932	8	7	0.7826
ق بِج CUL1	31	20	0.0192	11	3	0.0203
G IGF2BP3	25	19	0.168	3	2	0.6699
RELB	25	17	0.0741	3	0	0.1079
SEMA3E	19	7	0.006	1	0	0.4811

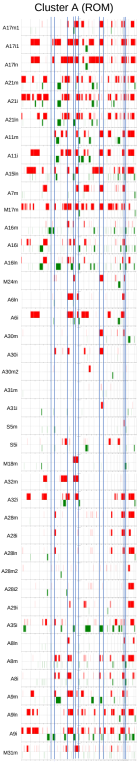
p values <0.05 are shown in bold. N/A: not assessable.

Online suppl. Fig. 1

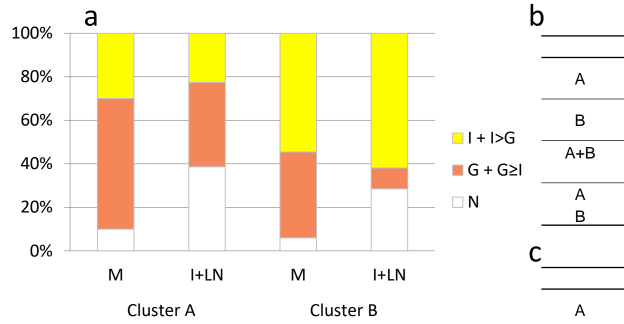




M34m



Online suppl. Fig. 2



			G + G≥l	Ν	p value
	А	Μ	12	2	0.0392
	A	I+LN	12	12	0.0392
	В	Μ	13	2	0.0062
	D	I+LN	2	6	0.0002
A	∖+ Β	Μ	25	4	0.0011
		I+LN	14	18	
	А		24	14	1.0000
	В		15	8	
			l + I>G	Non-I	<i>p</i> value
	А	Μ	6	14	0.7432
	Λ	I+LN	7	24	0.7452
	В	Μ	18	15	0.7784
		I+LN	13	8	0.7704
ŀ	∖+ Β	Μ	24	29	0.5545
		I+LN	20	32	
	A		13	38	0.0014
	В		31	23	

						Mean	T/R*	Difference (A vs B)	
No	Gene symbol	Gene name	Gene function	Location	Probe number	Cluster A	Cluster B	p value	p value after BC¶
1	VRK2	vaccinia related kinase 2	apoptosis and growth	2p16.1	4	0.2216578	-0.091428	1.78882E-06	0.026791191
2	XIRP2	xin actin binding repeat containing 2	actin binding	2q24.3	8	0.2525296	0.0062581	1.27706E-06	0.019126541
3	CNTN6	contactin 6	cell adhesion	3p26.3	6	0.00099	-0.322622	8.6214E-07	0.012912274
4	CNTN4	contactin 4	axon connections	3p26.3-p26.2	19	0.1137656	-0.189603	2.02011E-09	3.02551E-05
5	RBMS3	RNA binding motif single stranded interacting protein 3	c-myc gene binding	3p24.1	16	0.0883685	-0.259709	1.30358E-12	1.95238E-08
6	ARPP-21	cAMP regulated phosphoprotein 21	nerve function	3p22.3	5	0.1407465	-0.211016	3.69533E-07	0.005534503
7	STAC	SH3 and cysteine rich domain	SH3 and cysteine rich	3p22.3-p22.2	4	0.0944739	-0.24017	5.63687E-07	0.008442343
8	MYRIP	myosin VIIA and Rab interacting protein	myosin interacting	3p22.1	10	0.1020163	-0.178061	1.24906E-06	0.018707211
9	CACNA2D3	calcium voltage-gated channel auxiliary subunit alpha2delta	tumor suppressor gene	3p21.1-p14.3	20	0.1946772	-0.121071	3.18111E-08	0.000476436
10	PTPRG	protein tyrosine phosphatase, receptor type G	tumor suppressor gene	3p14.2	14	0.0442553	-0.238805	3.32493E-06	0.049797535
11	LRIG1	leucine rich repeats and immunoglobulin like domains 1	tumor suppressor gene	3p14.1	3	0.2577084	-0.234233	1.39033E-06	0.020822901
12	SLIT2	slit guidance ligand 2	tumor suppressor gene	4p15.31	11	0.0410187	-0.214485	1.32879E-06	0.019901247
13	SEL1L3	SEL1L family member 3	lymph node/stomach expression	4p15.2	3	-		5.63456E-07	0.00843888
14	FSTL5	follistatin like 5	tumor suppressor gene	4q32.2	16	0.0087773	-0.252876	1.05674E-06	0.015826866
15	GPM6A	glycoprotein M6A	oncogenic potential gene	4q34.2	9	0.1142247	-0.225603	2.75025E-07	0.004119045
16	MCCC2	methylcrotonoyl-CoA carboxylase 2	carboxylase	5q13.2	3	0.1370447	-0.323169	2.31579E-07	0.003468354
17	NXPH1	neurexophilin 1	nerve function	7p21.3	6	0.4400862	0.0622393	6.47209E-07	0.009693249
18	H2AFV	H2A histone family member V	histones nucleosome	7p13	2	0.6194794	0.1092316	8.29559E-07	0.012424304
19	SUN3	Sad1 and UNC84 domain containing 3	testis expression	7p12.3	2	0.3635659	-0.120918	8.55676E-07	0.012815461
20	COBL	cordon-bleu WH2 repeat protein	actin regulator	7p12.1	8	0.4966139	0.111728	4.98461E-08	0.000746545
21	SEMA3E	semaphorin 3E	proto-oncogene?	7q21.11	7	0.2570146	-0.078017	1.71778E-06	0.025727224
22	MUC12	mucin 12, cell surface associated	tumor suppressor gene	7q22.1	2	0.7944379		8.58551E-08	0.001285852
23	ORAI2	ORAI calcium release-activated calcium modulator 2	calcium modulator	7q22.1	2	0.370068	-0.099438	3.29493E-07	0.004934813
24	PRKD1	protein kinase D1	target for oncogenic KRas signaling	14q12	9	0.0830602	-0.212465	1.15426E-06	0.017287292
25	SLC39A9	solute carrier family 39 member 9	proto-oncogene	14q24.1	3	0.2246334	-0.265776	2.34539E-08	0.000351268
26	NRXN3	neurexin 3	nerve function	14q24.3-q31.	1 29	-0.009645	-0.199779	1.05375E-06	0.015782062
27	CHST14	carbohydrate sulfotransferase 14	sulfotransferases	15q15.1	1	0.5904759	-0.107594	2.16714E-06	0.032457219
28	HS3ST3A1	heparan sulfate-glucosamine 3-sulfotransferase 3A1	tumor growth factor related gene	17p12	5	0.1581726	-0.186387	2.87768E-06	0.043099021
29	MAP2K7		tumor growth factor related gene	19p13.2	2	0.3651435	-0.067787	3.70778E-07	0.005553136
30	SCUBE1	signal peptide, CUB domain and EGF like domain containing 1	EGF (epidermal growth factor)-like	22q13.2	4	0.0642863	-0.256986	5.19618E-07	0.007782325
31	MPPED1	metallophosphoesterase domain containing 1	brain/liver expression	22q13.2	3	0.482682	-0.047041	1.24649E-07	0.001866868
32	CELSR1	cadherin EGF LAG seven-pass G-type receptor 1	non-classic-type cadherin	22q13.31	4	0.0170639	-0.325309	2.06694E-06	0.030956486

*T/R: Tumor/Reference fluorescence intensity ratio. **¶BC:** Bonferroni corretion.