の段階でリンパ節転移を来たし、全身疾患としての性格 をもつことから、化学放射線療法については、局所治療 としての放射線療法と全身治療としての化学療法という 理解の上、理にかなった治療戦略として推進されている?

1999 年、Ohtsu らにより提唱された FP+RT 療法"は、その良好な成績から、非常に大きな注目を集め、多数例による長期間の予後に関する情報が収集された^{4,3)}。また、FP+RT療法と同様のプロトコールが欧米でも試験され、FP+RT療法と同様に、良好な結果が得られることが明らかにされた^{10,11)}。わが国では、さらに、UICC Stage II / III (T4を除く)の胸部食道扁平上皮がんに対するJCOG 9906 の試験が進行中であり(2000.4~2002.3 に 76例の登録を完了、現在症例追跡中)、中間解析ではあるが、37 例中 CR 24 例(65%)という良好な結果が得られている「コ、以上のような経緯を経て、現在、わが国では、FP+RT療法が、食道が入、特に進行性の食道がんに対する標準的治療法と認識されている。

本研究では、当院にて FP+RT 療法を受けた患者を対象に 2 年間の予後を追跡調査し、病期あるいは FP+RT 療法による治療終了約 4 週間後における評価と、予後との関係を解析した. 当院における奏効率は 80% であり、Kaneko 6²⁰の 81%、Ohtsu 6¹⁰の 87% とほぼ同程度であった。また、Stage I / II / III における奏効率が 80%以上に達しており、FP+RT療法が進行食道がんに対しても有効であることが確認できた。一方、2 年生存率は全体で 52% であった。Ohtsu 6¹¹の 1 年生存率 41%、3 年生存率 23%、Tahara 6⁴、もしくは、Ishikura 6²⁰の 3 年生存率 37-38%、また、スケジュールは同じであるが、CDDP や 5-FU の総投与量、放射線の総線量ともに、FP+RT療法より多い JCOG 9516¹³¹での 2 年生存率 31.5% と比較して、同等、あるいはやや優れている結果であった。

わが国における FP+RT療法に関する報告は4.50の多く は、主にT3/4を対象としており、病期と予後の関係を 明確にすることは難しい. 当院では、T1/2に対しても、 T3/4とほぼ同等の比率で、FP+RT療法を適用されて おり、その結果、図2に示すように、病期と予後が相関 することが明らかになった。また、図3に示すように、 治療終了約4週間後における評価で、CR 例と比較し て、non-CR 例において生存率が著しく低下することも 明らかとなった、ところで、Tahara らの報告の、もしく は Ishikura らの報告がは、FP+RT 療法に関する代表的な 報告であると位置づけることが可能である. 彼らの報 告4.57では、T 1/2 率 が 19%、Stage I / II 率 が 31% の 患 者に対して, CR率が56-58%, 2年生存率が40-45%, CR 例の2年生存率が70-75%, となっており、一方、 本研究では、T1/2率が43%、Stage I/II率が41%に 対して、CR率が45%、2年生存率が52%、CR例の2 年生存率が88%、となった、両者の比較から、当院では、非進行性の食道がんが多く、CR率はやや低く、予後は同等、あるいはやや優れている、という結果であると判別された、非進行性の食道がんが、進行性のものと比較して、良好な結果が得られることから、当院ではCRと判定されがたい、と考えられ、このことは、当院で、CR例の2年生存率が高いことと矛盾しないと思われた。当院では、治療終了約4週間後の評価で、1cm未満のリンパ節の腫脹が認められる場合、常法に従い、"uncertain CR"と評価し"、さらに、3カ月後に再検査を行う。このことがCR判定港準の違いに少なからず関与しているもの、と推察された"。

本研究では、T因子、M因子によっても予後が推定できる可能性が示された。ところで、Kaneko ら³¹は、T因子により予後の推定が可能であるものの、M因子については推定が困難であることを示している。具体的には、T3とT4の生存期間(中央値)はおのおの29カ月と11カ月であり、統計学的に有意な差異が認められるものの、M0とM1ではおのおの15カ月と12カ月であり、大きな違いがないことを報告している。T因子、M因子による予後の推定についてはさらに詳細な検討が必要であると思われた。なお、2年生存率は、5-FU投与量(mg/day)に依存しないものと推察された。

以上,2年間の予後を追跡調査し、病期あるいは治療終了約4週間後における評価と、予後との関係を解析した。その結果、1)当院における奏効率は80%であり、わが国における他施設の結果と同等であること、2)進行性の食道がんに対しても有効であること、3)2年生存率は52%であり、わが国における他施設の結果と同等、あるいはやや優れていること、4)病期と予後が相関すること、5)治療終了約4週間後における評価、具体的には著効したか否か、で予後が決まること、6)T因子、M因子も予後の推定因子になり得るかもしれないこと、を確認もしくは明らかにした。当院におけるFP+RT療法の治療成績も他施設の結果と同等であり、FP+RT療法は、施設に依存することなく高い治療成績を得ることができることから、食道がんにおける標準治療法となる可能性が示唆された。

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Research Paper

Analysis of p53 mutation status in human cancer cell lines

A paradigm for cell line cross-contamination

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Key words: p53 mutations, cancer, human cancer cell lines, NCI-60 panel, cell lines cross contaminations

Cancer cell lines are essential tools used in many areas of biomedical research. Using the last release of the UMD_p53 database (2007) (http://p53.free.fr), we analysed the p53 status of 1,211 cell lines published between 1989 and 2007, p53 mutations in cell lines from various types of cancers display a striking similarity in the distribution of mutations and in the pattern of mutational events compared to tumors, indicating that they are representative of the cells from which they were derived. Analysis of the residual transcriptional activity of p53 mutants identified in cell lines that displayed two different p53 mutations show that there is a high frequency of weak mutations that are paired with more potent mutations suggesting a driver/passenger configuration. Strikingly, we found discrepancies in the p53 status for 23% (88/384) of cell lines, for which the p53 status was established independently in two laboratories. Using the NCI-60 cell line panel as a model widely used in the literature, the p53 status could not be ascertained for 13 cell lines due to a lack of homogeneous data in the literature. Our study clearly confirms that misidentified cell lines are still a silent and neglected danger and that extreme care should be taken as a wrong p53 status could lead to disastrous experimental interpretations. The p53 web site has been updated with new sections describing the p53 status in the majority of cell lines and a special section devoted to cell lines with controversial p53 status.

Introduction

Continuous cell lines derived from human tumors are widely used in laboratory research. They can be used for drug screening (the NCI-60 panels), for production of various macromolecules, for modelling human tumors or, most frequently, as biological test tubes for a large variety of experiments. To draw valid conclusions from such experiments, it is essential for cell lines to be clearly characterized at the molecular level. For a long time, these genetic characterizations

were performed by studies focusing on one gene and the information was scattered in the literature. Recently, the Sanger Institute developed a Catalog Of Somatic Mutations In Cancer (COSMIC) that gathers information on genetic alterations in human tumor cell lines.² To date, data in the COSMIC cell line database is a mix of information taken from the literature and in-house sequencing.^{2,3}

Cell line cross-contamination (CLCC) is not a novel problem. ^{4.5} as it was discovered as early as 1974 that one in three cell lines were contaminated, mostly by HeLa cells. ⁶ Despite the tremendous work conducted by Nelson-Rees et al., this problem is still "a silent and neglected danger", as a recent study indicates a CLCC of 18% at a German cell line repository. ^{4.7,8} CLCC is not trivial, as the use of wrong cell lines can lead to erroneous conclusions associated with years of wasted time and effort. ⁹⁻¹¹

p53 mutation is the most common genetic abnormality found in human cancer. ¹² In cell lines, loss of p53 activity is usually linked with several specific landmarks such as defect in growth arrest or apoptosis after DNA damage and lack of induction of p53-regulated genes. ^{13,14} The p53 status is also a key factor for the sensitivity to anticancer agents and multiple studies have focused on this subject. ^{15,16} Although the majority of studies found a correlation between loss of p53 function and p53 alteration, a few publications report opposite results. ^{15,17,18} This situation is complicated by the observation that some mutant p53 proteins expressed in cell lines have only a partial loss of activity or present a temperature-sensitive transcriptional activity. ¹⁹

For more than 17 years, we have collected and compiled p53 mutations in human tumors and cell lines. 20,21 Although numerous studies on p53 mutations in human tumors have been published, no systematic analysis of the p53 status of cell lines is currently available. In the course of updating the various versions of the UMD p53 database, we have noticed a number of discrepancies in the p53 status of several cell lines. The situation has recently been worsened, as these discrepancies have been randomly published in the literature, a situation that can lead to serious problems of data analysis. Many drug sensitivity studies are based on the p53 status reported in the literature without any new genetic analysis.

In the present study, using the UMD-p53 database as a framework, we performed a precise and thorough analysis of p53 status in 1,211 tumor cell lines. Our analysis shows that p53 mutations in cell lines from various types of cancers display a striking similarity in

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the distribution of mutations and in the pattern of mutational events when compared to tumors indicating that they are representative of the cells from which they were derived. Surprisingly, we found discrepancies in the p53 status in 23% of cell lines, some of which are widely used, such as MOLT-4 or CAPAN-1.

Results and Discussion

p53 mutations in cell lines versus tumors. The pattern of p53 mutations can be analysed in two informative ways, either by examining the distribution of p53 mutations in the p53 protein or by scoring the various mutational events that lead to these mutations. Both types of analysis have been very informative when applied to various types of human tumors.²⁵ These studies demonstrate a link between exposure to various types of carcinogens and the development of specific cancers. The most striking example is that of tandem mutations, specifically induced by ultraviolet radiation, which are only observed in skin cancers. 26 The relationships between G-T transversion and lung cancer in smokers or mutation of codon 249 observed in aflatoxin B1-induced liver cancers are also very demonstrative. 25,27 The distribution of p53 mutations along the p53 protein is similar in tumors and cell lines, indicating that there is no bias in the selection of specific mutant p53 during establishment of a cell culture (Fig. 1A and data not shown). The only exception concerns colorectal cancer cell lines. p.R175H is one of the most frequent p53 mutations in tumors, but is very rare in colorectal cancer cell lines (Suppl. Fig.). This finding is specific for p.R175H and it is not observed for the other two hot spots at codons 248 and 273. The reason for this bias is not known. Comparison of the various mutational events in cell lines and tumors has been performed for all cancers together or for 8 cancer types (Fig. 1A and B and Suppl. Figs.). As previously observed, the pattern of mutations differs between various types of cancers, but there is a striking similarity when comparing tumors and cell lines from the same origin. In colorectal and brain cancer, there is a predominance of GC-AT transition at CpG dinucleotides, whereas in lung cancer or head and neck SCC, the frequency of GC-TA transversion is 30% and 20%, respectively, with only a few transitions at CpG dinucleotides. This high frequency of transversion in these cancers has been shown to be associated with tobacco smoking and will not be discussed in more detail here.²⁸ This similarity in the pattern of p53 mutations in primary tumors and cell lines is a strong argument suggesting that these p53 mutations did not occur de novo during the establishment of these cell lines. It also supports the small number of studies that have found matched p53 mutations in primary tumors that were used to establish cell lines and confirms that analysis of the spectrum of mutations in oncogenes or tumor suppressor genes in human cell lines accurately reflects the situation observed in primary tumors.

Analysis of p53 mutant activity in cell lines. Analysis of p53 mutations in human tumors has led to the discovery that at least 5% to 10% of published p53 mutations could be due to PCR or sequencing artefacts. ²² However, these mutations are not randomly distributed among the 2,500 publications reporting p53 mutations. A meta-analysis identified about 30 publications (1,600 p53 mutations) with a high concentration of unusual p53 mutations that shared the following properties: (i) multiple p53 mutations in the same tumor (3 to 14); (ii) a high frequency of synonymous mutations; (iii) a low frequency of mutations at hot spot codons; (iv) most of these

mutations retained either partial or total transactivational activity.²⁹
The vast majority of these studies were associated with the use of nested PCR for amplification and analysis of the p53 gene. Analysis of p53 mutations in cell lines provides several advantages over analysis of tumors to minimize artefactual data: (i) DNA extracted from cell lines is available in large quantities. Analysis requires neither nested PCR nor excessive numbers of PCR cycles and can be easily repeated; (ii) The high quality of the DNA avoids PCR problems associated with DNA extracted from paraffin-embedded tissue; (iii) DNA is not contaminated by normal DNA from stroma or cells or infiltrating lymphocytes.

The UMD p53 mutation database includes functional information about the majority of p53 missense mutants, as originally published by Kato et al.,23 (see also material and methods). Quantitative data concerning the transcriptional activity of each missense p53 mutation has been extremely useful to classify and analyse p53 mutations in the p53 database.21,22,29 The mean and 95% confidence interval (CI) of the remaining activity of all mutant p53 proteins found in cell lines or in tumors was calculated by using the activity measured on the p21WAF1 promoter (similar results were obtained with the activity measured on seven other promoters of transcription, data not shown). The analysis shows that the mean activity was situated between -1 and -1.2. This value corresponds to a residual transcriptional activity of about 10% compared to wild-type p53. The narrower 95% CI in tumors compared to cell lines is due to the greater number of tumors used in the analysis (Fig. 2A). In the majority of cancers, residual p53 activity was lower in cell lines than in tumors, but this difference was only marginally significant in head and neck, breast and SCLC, p = 0.03). On the other hand, residual p53 activity has a wider distribution in tumors compared to cell lines (variance analysis, Fig. 2B). A large number of mutant p53 retain wild-type activity in tumors, but this feature is rarely observed in cell lines. This difference was highly significant for all cancer types (p < 0.0001) except for brain cancers and haematological malignancies. Two non-exclusive explanations can be proposed for this difference between tumors and cell lines. First, it is possible that only tumors with fully inactivated p53 are preferentially selected to establish cell lines. This hypothesis could also explain why the frequency of p53 mutations is always higher in cell lines than in tumors. It is also possible that this profile of p53 inactivation in cell lines is more representative of the true pattern of p53 inactivation and that the tumor p53 database contains passenger mutations and/or artefactual mutations with partial or fully active p53.21

During the course of these analyses, we also observed that 82 cell lines displayed two p53 missenses mutations. Preliminary observations suggested that the two mutations may not have the same importance and that only one mutation was the driving force selected during transformation. ²² In order to obtain more information, clustering analysis was performed on cell lines with either single (SM cell lines) or double mutations (DM cell lines). Three clusters were obtained for the two populations, corresponding to mutant p53 with wild-type activity (cluster II) intermediate residual activity (cluster II) or no activity (cluster III) (Table 1). The number of mutants in clusters I and II was significantly higher in DM cell lines than in SM cell lines, whereas mutations with total loss of activity were more frequent in SM cell lines (p < 0.0001, Table 1). Mutations in DM cell lines were further analysed to determine how paired mutations

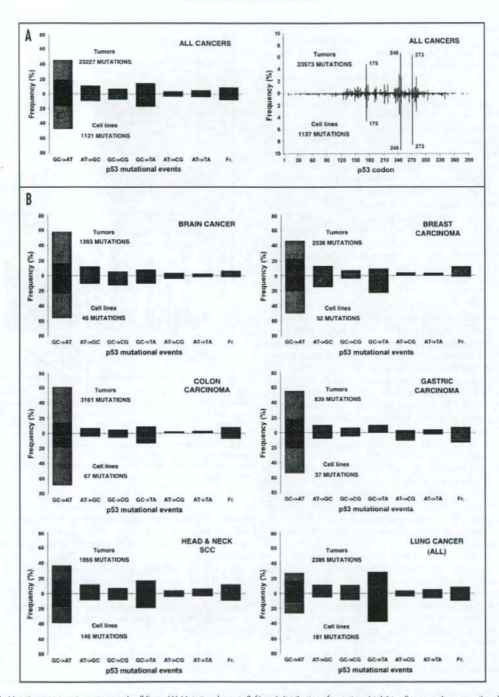


Figure 1. Mutation spectrum in tumours and cell lines: (A) Mutational events (left) and distribution of mutations (right) in all tumours (upper part) and cell lines (lower part). Data were obtained from the UMD p53 database, 2007_R1 release (http://p53/free/fr]. (B) mutational events in tumours versus cell lines in various types of cancer. A similar pattern of mutational events is observed for other cancers (melanoma, ovarian carcinoma, oesophageal carcinoma or pancreatic carcinoma, data not shown). Transitions at CpG dinucleotides are shown in red.

(two mutations in a single cell line) were associated (Table 2). Only one of the 41 cell lines presented two mutations in cluster I with wt activity for the two p53 mutant alleles. This choriocarcinoma cell line (NUC-1) displays two unusual p53 mutations at codons 17 and 24 that have never been observed in any other cell lines or tumors. Among the 11 remaining mutations in cluster I, three were paired with mutations in cluster II and 8 were paired with mutations in cluster III. Among the 19 mutations in cluster II, two were paired with a mutation of the same class, 3 with class I mutations and 12 with class III mutations. The majority (30) of the 50 mutations in cluster III were paired with a mutation of the same class and 12 were paired with class II mutations (Table 2).

Double mutations can occur in two configurations, either on the same allele (DMS) or on two different alleles (DMD). Unfortunately, in the majority of cases, this status is unknown (DMU). In the p53 mutation database based on tumors, the majority of DM with a known configuration are DMD (about 90%). No cell lines with two missense mutations in the same allele have been reported and only 10 cell lines with mutations on two different alleles have been reported. All of these cell lines expressed one class III mutation associated with either another class III mutation (6), or class II (3) or class I (1) mutations.

Altogether, our results indicate that: (i) there is a higher frequency of weak mutations in DM than in SM mutations and (ii) the majority of these weak mutations are paired with a more potent mutation. This suggests that the two mutants do not have the same contribution to the transforming process. Whether or not these weak mutations are passenger mutations associated with a driving mutation or true mutations associated with selection of the transforming phenotype is an unresolved question. One of the main problems associated with p53 mutations is the possible dominant negative activity of mutant p53 via hetero-oligomerization making it very difficult to reach any definitive conclusions concerning weak p53 mutations. Weakening of the second allele could possibly accentuate the dominant negative activity of p53.

p53 status in human tumor cell lines. The NCI-60 panel is a good example of a series of cell lines that are widely used for both basic research and drug discovery.1 This panel originally contained 60 cell lines from nine histological origins (Table 3). Several observations unrelated to p53 status revealed that some cell lines were either mixed up or were derived from the same donor (Table 3).5 At least 100 studies have analysed the p53 status of a subset of the panel and in 1997, O'Connor et al., reported the p53 status of the entire NCI-60 panel. 16 This paper has been used as a reference for 10 years despite discrepancies with other data in the literature. A second analysis of the entire NCI-60 panel was performed in 2006 and the results are fairly heterogeneous compared to the 1997 study (Table 3). Inspection of the two studies leads to the detection of 19 apparent differences (Table 3). Three differences were due to typographical errors in the 1997 report (RPMI-8226, SK-MEL-28 and Hs-578-T). A more careful examination of four other discrepancies reveals that they are due to a problem of nomenclature associated with a different mutation screening strategy. In the 1997 paper, p53 mutations were analysed by cDNA sequencing, while the 2007 analysis was performed using genomic DNA as starting material. One of the disadvantages of RNA-based analysis is that it is impossible to infer whether deletions found in the cDNA are due to splicing

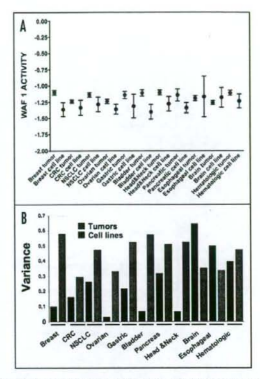


Figure 2. Analysis of the residual p53 activity of mutant p53 in tumors and cell lines. (A) Points, mean p53 activity as measured by transactivation with the p21WAF1 promoter; bars, 95% Cl. A similar distribution was observed with other p53 response genes (data not shown). The yaxis corresponds to p53 transactivation activity, with a value of -1.5 for the negative control and 2.5 for wild-type p53. (B) Variance of the p21WAF1 promoter activity in tumors and cell lines. CRC, colorectal carcinoma; NSCLC, non-small cell lung cancer. Data from cell lines and tumors are displayed in black and red respectively.

mutations or intragenic deletions in the gene. On the other hand, it is always difficult to predict the consequence of mutations found in intron or splice junctions after genomic sequencing. Both methods are complementary and may be necessary to ensure an accurate genetic status.

In HOP-62, RNA-based analysis detected an insertion between codon 212–225 but no information about the insertion sequence was available. Codon 225 is at the boundary of exon 6 and intron 6 suggesting a splicing defect, as analysis at the genomic level confirms the presence of a splice mutation in the acceptor signal of exon 6 (Table 3).

In OVCAR-8, the 126–132 deletion detected by the RNA-based assay concerns the first six residues of exon 5. Genomic analysis described a mutation in the acceptor site of exon 5 and a splicing defect leading to a shift of the normal donor site of exon 5 that skips 18 nucleotides (6 aa residues) in exon 5. Examination of the DNA sequence at codon 132 reveals an AG dinucleotide sequence preceded by a pyrimidine tract similar to those found in the splice donor sequence. The same situation is observed for NCI/ADR-RES

2008; Vol. 7 Issue 5

Table 1 Cluster analysis of p53 mutation activity

	SM	DM
Cluster I (wt activity)	27 (3.4%)	13 (15.9%)
Cluster II (low acitivity)	73 (9.3%)	19 (23.2%)
Cluster III (no activity)	687 (87.5%)	50 (61.0%)
Total	787 (100%)	82 (100%)

The table entries are the number (and %) of mutants dassified into the three dusters based on k-means dustering of the promoter activities of pS3 target genes. There are significantly more duster-1 and duster-11 mutations among the double mutations (DM) than among the single mutations (SM) (p = 2e - 10 using the hi-square test).

Table 2 Discordance table of class assignment of the 82 DM mutations (from 41 pairs)

	Cluster I	Cluster II	Cluster III
Cluster I (wt activity)	1	3	8
Cluster II (low acitivity)	0	2	9
Cluster III (no activity)	2	3	15

Majority of the weak mutations (duster I and duster II) are paired with strong mutations (duster III).

that has been recently shown to be an ovarian carcinoma cell line originating from the same patient as OVCAR-8.

In EKVX, the deletion of codon 187 to 224 detected on RNA-based analysis corresponds exactly to the deletion of the entire exon 6, a strong argument for a splicing defect. Genomic analysis did not reveal a splicing defect but a tandem mutation at codons 203 and 204 in exon 6 (Table 3). If the two cell lines analysed were really EKVX, this result suggests that a mutation at either codon 203 and/or 204 could affect p53 gene splicing. This observation is not surprising, as it is now well known that exons contain exonic splicing enhancers (ESE) that regulate either alternative splicing or normal splicing.30 These ESE are recognized by the SR proteins that regulate the various splicing events. Mutations in ESE have been identified in numerous genes including APC or NF1.31,32 Exonic mutations that can change p53 splicing have also been described. 33,34 Taken together, the contradictions noted in the p53 status of the four cell lines, HOP-62, OVCAR-8, NCI/ADR-RES and certainly EKVX are only due to the different strategies used for their analysis and a lack of homogeneity in the nomenclature used to report p53 mutations. The problem of the nomenclature of p53 mutations as well as other gene defects is a recurrent problem in publications.³⁵ Despite numerous recommendations, the description of p53 mutations in the literature is highly heterogeneous and can reach a high degree of fantasy with tables that are either totally non-informative or with so many typographical errors that they cannot be interpreted. In a recent survey, the editors of 80 journals with frequent publications of p53 mutations were contacted in order to stress this problem and define certain guidelines for the publication of p53 mutations (Soussi T, Unpublished). Unfortunately, this survey was a complete failure with less than 10% of replies and no change in the trends of reporting accurate p53 mutations. In fact, the number of typographical errors or incomprehensible mutations has increased over the last five years (Soussi T, unpublished observations).

After eliminating typographical errors and possible splice mutations, the p53 status of 15 cell lines was different between the two studies. Using the UMD p53 database and the literature, we checked for other publications that have analysed the p53 status of these cell lines. For two cell lines, CCRF-CEM and HL-60, sufficiently concordant publications are available to define a consensus concerning the p53 status (Table 3). For 13 cell lines, analysis of the literature revealed a very heterogeneous situation and no consensus could be reached (Table 3, Inconclusive). Cell lines such as MOLT-4 or NCI-H226 represent an extreme situation, as multiple publications do not show any common p53 mutations. For other cell lines such as DU-145, which have been shown to display two different p53 mutations in two different alleles (p.V274F and p.P223L), the ambiguity concerns the fact that several authors have detected only one of the two mutations, either p.V274F or p.P223L. It is therefore possible that during long-term cell culture, one of the two mutant p53 alleles is lost, as no selection pressure is exerted on cell growth.

A similar situation is observed for other cell lines that do not belong to the NCI-60 panel, but with many discrepancies (Table 4 and Suppl. Table S1, see also p53 website). In many cell lines, the p53 status has been analysed in only one or two reports and the information is subsequently reproduced in the literature. This is a very dangerous situation as it could lead to erroneous phenotypegenotype correlations in various types of studies. The pancreatic carcinoma cell line CAPAN-2 is a good example of the problems raised by erroneous phenotype. This cell line has been described as either wt, mutated (p.R273H) or p53 null (Table 4 and reference within). A Pubmed literature search indicates that all three phenotypes are used in various studies.

The "p53-null" status is used in different ways in the literature. The two most common meanings are a cell line with a documented p53 gene deletion (both alleles) or a cell line with a p53 mutation. We have also observed more "unusual" situations in which this status is only based on p53 expression (RNA or protein). Unfortunately, this type of information diffuses rapidly in the literature without any verification of the original publication. The p53 status of the two cell lines SK-OV-3 (Ovarian cancer) and FRO (anaplastic thyroid carcinoma cell line) are a good example of this ambiguity. In the majority of publications, the p53 status of these two cell line is stated as "p53 null". In fact, close examination of the original manuscript shows that the p53 gene in SK-OV-3 is not deleted and did not sustain any gross rearrangement but neither p53 RNA or protein are found. In these publications, no p53 murations were found but the recent analysis performed at the Sanger Institute detected a deletion of a single nucleotide at position 267 (codon 90).3 It is therefore possible that nonsense-mediated mRNA decay (NMD) eliminates p53 aberrant mRNA. NMD has been observed in the human leukaemia cell line K562 where p53 is also inactivated via a 1 base pair insertion at nucleotide 136. For the FRO cell line, the original reference for the analysis of the p53 gene status is always correctly quoted, but a closer look at this original paper demonstrates a marked decrease of p53 RNA in the cell but no mutation was detected by sequencing exons 5 to 8. Either a mutation is situated outside this region leading to a decrease of RNA expression (frameshift mutation associated with Nonsense-Mediated mRNA Decay) or the altered p53 expression is due to another mechanism. Because the whole p53 gene is present, it is incorrect to define SK-OV-3 or FRO cell lines as "p53 null", as

Table 3 p53 status in the NCI-60 panel cell lines

Cell line	ATCC	Cancer	Mutation in UMD ¹		Mutation in COSMIC ²		Consensus ²	Comments	
	number		Mutation	Reference	Protein	DNA			
CORF-CEM	CCL-119	Leukemia	R248Q	16	p.R175H, p.R248Q		p.[R175H] + p.[R248Q]	Two mutations in separate alleles. Derivative CCRF-CEM-	
			R175H+ R248Q	394			pinzaouj	VLB100 has a third mutation	
HL-60 ⁸	CCL-240		R248L	16	p.M1_*394del	c.1_1182del1182	p.M1_*394del		
			p53 Null	40 4					
K-562	CCL-243		ND	16	p.Q136fs*13	c.406_407insC	p.Q136fs*13	W. State Street Street	
	LIGHT IN	100	136ins1	41 4			1 11 11 11 11	And the second	
MOLT-4	CRL-1582		wt	16,39	p.R306X	c.916C>T	Inconclusive		
			L111V	42				1 14 2 1	
			R248Q	43					
RPMI-8226	CCL-155		E285L'	16	p.E285K	c.853G>A	p.E285K	This mutant is temperature-	
			E285K	44				sensitive	
SR	CRL-2262		wt	16	wt.	wt	wt		
A549	CCL-185	Non-Small Cell Lung	wt	16,45	wt	wt	wt	Dalo al la	
EKVX			del187-224	16	p.V203_E204>V*	c.609_610GG>TT	Inconclusive	Possible splicing defect See text for discussion	
HOP-62			Ins212-225	16	p.?	c.673-2A>G	Splicing defect	See text for discussion	
HOP-92	-		R175L	16	p.R175L	c.524G>T	p.R175L		
NCI-H226	CRL-5826	77 77	P309A	16	wt	wt	Inconclusive		
		C 1942	R158L	46		300			
NGI-H23	CRL-5800		M246I	16,47	p.M246I	c.738G>C	p.M246l		
NCI-H322M	255-25-57		R248L	16.46	p.R248L	c.743G>T	p.R248L	Martin Street	
NCI-H460	HTB-177		wt	16	wt	wt	wt		
NCI-H522	CRL-5810	2 37 7 7 7	191delG	16 46	p.P191fs*56	c.572_572delC	p.P191fs*56	THE REAL PROPERTY.	
COLO-205	CCL-222	Colon	G266E	16	p.Y103_L111>L	c.308_333>TA	Inconclusive		
	-		103del27	48					
HCC-2998	1		R213X	16	p.R213x	c.637C>T	p.R213x	Taran and Inc.	
HCT-116	CCL-247		wt	16	wt	wt	wt		
HCT-15	CCL-225	1911316	P153A	16	p.S241F, p.?	c.722C>T, c.1101-	Inconclusive	HCT-15 and DLD1 are derived	
	100000		S241F	49		2A>C		from the same individual. DLI display a p.S241F mutation	
HT-29	HTB-38		R273H	16 48	p.R273H	c.818G>A	p.R273H		
KM12	1	O MITTER	H179R	16	p.R72fs*51	c.215delG	Inconclusive	CONTRACTOR OF THE PARTY	
SW620	CCL-227		R273H	16 50	p.R273H, p.P309S	c.818G>A, c.925C>T	p.[R273H] (+)	SW480 and SW620 are derived	
			R273H, P309S	51			p.(P309S)	from the same individual with a similar p53 atteration. The p.P309S mutation is not always reported	
SF268	1	CNS	R273H	16,52	p.R273H	c.818G>A	p.R273H	THE REAL PROPERTY.	
SF295			R248Q	16	p.R248Q	c.743G>A	p.R248Q		
SF539	TEN IN	100	wt	16	p.R342fs*3	c.1024delC	Inconclusive	MENN DEPT TOP	
SNB75	/ River in	50 311	E258K	16	p.E258K	c.772G>A	p.E258K		
U251/SNB19			R273H	16,53	p.R273H	c.818G>A	p.R273H	SNB19 and U251 cell lines are derived from the same individual and are similar	

in the case of H1299 or Saos-2 cell lines in which the p53 gene is entirely deleted. These cell lines are commonly used as recipients to reintroduce either wild-type of mutant p53. Whether the presence of an endogenous p53 gene which is still transcriptionally active in the SK-OV-3 or FRO cell could interfere with this reconstitution experiment is not known, but should be carefully considered before conducting this type of experiment. The recent finding of p53 isoforms that could be expressed by alternative splicing may also increase the complexity of this problem, as the various delta133 isoforms could be theoretically expressed in this cell line.

Another reason why "p53-null" should be used cautiously to describe cell lines that express mutant p53 is the observation that p53 mutations

are fairly heterogeneous in terms of loss of function and several cell lines display a normal or partial p53 response. Finally, there is now ample evidence that some mutant p53 behave as dominant oncogenes with a gain of function activity. We therefore believe that the "p53 null" status should be used only for cell lines that are totally devoid of p53 gene. Any other situation should be referred to as "mutant p53".

The UMD_p53 database (2007_R1 release) includes p53 mutations in 1,211 cell lines: 827 of these mutations have only been described once, preventing any verification. A discrepancy was detected in 88 of the remaining 384 cell lines (23%), in line with the study by Macleod et al., who showed that 18% of cell lines in the DSMZ-German Collection of Microorganisms and cell Cultures

2008; Vol. 7 Issue 5

Table 3 p53 status in the NCI-60 panel cell lines (continued)

LOXIMV1		Meianoma	wt	16	wt	wt	wt	
Makne-3M	HTB-64		wt	16	wt	wt	wt	- 11 - 1
M14	-		G266E	16	p.G266E	c.797G>A	p.G266E	DE LA COMPANIE DE LA
SK-MEL-2	HTB-68		G245S	16	p.G245S	c.733G>A	p.G245S	
SK-MEL-28	HTB-72		C145V'	16	p.L145R	c.434_435TG>GT	p.L145R	634
			L145R	54				- Contraction
5K-MEL-5	HTB-70		wt	16	wt	wt	wt	
UACC-257			wt	16	wt	wt	wt	
UACC-62			wt	16	wt	wt	wt	
MDA-MB-435			G266E	16, 55	p.G266E	c.797G>A	p.G286E	This cell line was originally reported as a breast carcinoma cell line but recent SNP analysi indicates that it is similar to the M14 melanoma cell line
MDA-N			G268E	16	ND	ND	p.G266E	This cell line is a derivative of MDA-MB-435 transfected with a plasmid expressing erbB2
IGROV1		Ovarien	wt	16 56	p.Y126C	c.377A>G	Inconclusive	
OVCAR-1	HTB-161		R248Q	16 57	p.R248Q	c.743G>A	p.R248Q	
OVCAR-4		7.0	wt	16	p.L130V	c.388C>G	Inconclusive	Carried Wallet
OVCAR-5			ins224	16	wt	wt	Inconclusive	
			224ins3*	58				
OVCAR-8	100		del 126-132	16	p.?	c.376-1G>A	Splicing defect	Same as NCI/ADR-RES
NCI/ADR-RES			del 126-132 126del21	16	p.?	c.376-1G>A	Splicing defect	Originally labelled as MCF- 7/Adr but was later found to be different from MCF-7. SNP analysis indicates that it is
								similar to OVCAR-8
SK-OV-3	HTB-77		H179R	16	p.S90fs*33	c.267delC	Inconclusive	
786-0	CRL-1932	Renal	P278A	16	p.P278A, p.?	c.832C>G, c.560- 2A>G	Inconclusive	
A498	HTB-44		wt	16	wf	wf	wt	
ACHN	CRL-1611	-	wt.	16	wt	wt	wt	
CAKI-1	HTB-46		wt	16	wt	wt	wt	
HXF393	-	1	R175H	16	p.R175H	c.524G>A	p.R175H	
SN12C			E336X	16	p.E336X	c.1006G>T	p.E336X	
TK10	7.00	1000	L264R	16	p.L264R	c.791T>G	p.L264R	
U031			wt	16	wt	wt	wt	
DU-145	HTB-81	Prostate	P223L	16	p.V274F	c.820G>T	Inconclusive	See text for more information
	FE.		P223L, V274F	60,61	E Samo	10000		
PG-3	CRL-1435		138del	16	p.K139fs*31	c.414delC	p.K139fs*31	
			138del1	60				
			R282W	62				
BT-649	HTB-122	122 Breast	ND 16 p.R249S	c.747G>C	p.R2495			
	100		R249S	63			The state of	in the second
Hs 578T	HTB-126		D157E'	16	p.V157F	c.469G>T	p.V157F	
			V157F	64.4				
MCF7	HTB-22		wt	16	wt	wt	wt	
MDA-MB-231	HTB-26		R280K	16, 63	p.R280K	c.839G>A	p.R280K	
T47D	HTB-133		L194F	16,51	p.L194F	c.580C>T	p.L194F	

Mutations as reported in the 2007_R1 of the UMD p53 mutation database. The description of the mutations have been left as originally published by the authors, ⁷Mutations described by Kediobi et al.,², ⁵A mutation consensus was defined for cell lines using the following rules: (i) at least two independent studies reporting sequencing and identifying the same mutation without any contradictory reports; (ii) at least three independent studies reporting sequencing and identifying the same mutation and one fourth contradictory report. All other possibilities were not considered to be consensual and have been assigned as uncertain. The namendature for TP53 mutation uses either the cDNA (RefSeqNM 000546.2) or the protein (RefSeq NP 000537) as reference. For numbering, +1 is A of the ATG initiation codon in the correct RefSeq (NM 000546.2). Mutations are described using the international nomenclature⁵⁵ and http://www.hgvs.org/mutnamen/; ⁴Mutation found independently by multiple authors. Only the first publication is shown; ⁵HL 60(TB) was used for the analysis, but it is reported to have a p53 deletion similar to HL60; "The status of MOLT-4 is highly heterogeneous in the literature. The report of a wt status could be due to the fact that only exons 5 to 8 (residues 126-306) were screened in several publications; "Typographical error in the publication; his not dear whether these authors checked the p53 status of the cell line or report the mutation described by 0' Connor et al., "This cell line has been reported to be null for pS3 RNA or protein. Whether this is due to a small DNA rearrangement or RNA-mediated decay associated with a frameshift mutation is unknown.

were cross-contaminated.8 The p53 status in various cell lines is a mutation hot spots (Ha-ras) or a lower frequency of mutations are not

paradigm for CLCC. (i) p53 mutation is sufficiently diverse to allow as useful. (ii) Due to its importance in cell phenotype, p53 status has comparison of various cell lines. Statuses of other genes with fewer been analysed in more than 1,200 cell lines. Although p53 mutation

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Table 4 Cell lines with controversial p53 mutations

Cell line	ATCC	Cancer	p53 Mutation UMD		
ACCOUNT OF THE PARTY OF THE PAR	number	***************************************	AA Change	Reference	
HT-1197	CRL-1473	Bladder carcinoma	p.His365Arg	66	
			wt	2	
EJ		Bladder carcinoma	p.Tyr126X	67	
			p.Lys164Glu	68	
			wt	69	
RT-112		Bladder carcinoma	p.Ser183X, p.Arg248Gln	2	
			p.Arg248Gln	70	
SD		Bladder carcinoma	p.Arg110Leu	71	
			p.Ser116Cys	67	
T-24	HTB-4	Bladder carcinoma	p.Tyr126X	67	
			p.Tyr126delTAC	66	
VM-CUB-1		Bladder carcinoma	p.Tyr126X, p.Arg175His	2	
			p.Arg175His	67	
VM-CUB-2		Bladder carcinoma	p.Arg158Leu, p.Tyr163Cys	67	
			p.Arg158Leu	71	
MDA-MB-436	HTB-130	Breast carcinoma	p.Glu204delinsAspfsX6	55	
	1.1.0		p.Arg273His	72	
DAUDI	CCL-213	Burkitt lymphoma	p.Arg213X	73	
DAGDI	0022.0	South Mary Transport	p.Gly266Glu	2	
A-172	CRL-1620	Glioblastoma	p.Cys242Phe	74	
	0112 1020		wt	75	
U-118-MG	HTB-15	Glioblastoma	p.Arg213Gln	76	
0-110-1110	11.0.10	- Citobiotoma	wt	2	
SK-LMS-1		Leiomyosarcoma	p.Met237Lys, p.Gly245Ser	77	
OIT EIIIO T		actionly obtained	p.Gly245Ser	78	
SK-UT-1	HTB-114	Leiomyosarcoma	p.Arg175His, p.Arg248Gln	2	
011-01-1	1110-114	Loioniyodardanid	p.Arg175His	78	
NCI-H1048	CRL-5853	Lung (SCLC)	p.Ser46fsX76, p.Arg273Cys	2	
1101-1110-10	CINE DOOD	Lung (DOLO)	p.Arg273Cys	79	
MeWo	HTB-65	Melanoma	p.Glu258Lys, p.Gln317X	2	
merro	1110-00	Wolanoma	p.Glu258Lys	80	
PA-1	CRL-1572	Ovarian carcinoma	p.Asn239Asp	81	
ra-i	JAC-1072	Ovarian CarcinOma	p.Pro316Pro	57	
2	1		p.Flo3foFlo wt	2	
SW626	HTB-78	Ovarian carcinoma	p.Gly262Val	58	
344020	H1B-78	Ovarian carcinoma	p.Giy202Vaii p.Arg273His	82	
Canan 2	LITE OC	Danomatic carres		83	
Capan-2	HTB-80	Pancreatic cancer	p.Arg273His wt	2	

^{*}References correspond to studies in which the p53 gene status was analysed experimentally and not deduced from other reports in the literature.

analysis cannot replace DNA fingerprinting, our finding is a strong argument to suggest that CLCC should not be ignored. We are also very concerned by the observation that the p53 status based on a cell line (either correct or false) can be reproduced from a single publication in the literature without any subsequent confirmation. Finally, we have also noticed a marked heterogeneity in the labelling of cell lines, a problem that can also lead to confusion between mislabelled cell lines with similar names. ³⁶ CLCC includes several situations: (i) cross-contamination between two cell lines (the best example being HeLa cells); (ii) cell lines with an incorrect origin (such as the KB cell line often wrongly described as an oral cancer when it is actually a cervical cancer); and (iii) cell lines that have been contaminated during manipulation. We believe that the problem identified in the

present analysis is predominantly related to confusion or incorrect labelling of cell lines. Although, the material and methods sections of published articles usually state that cell lines were derived from cell banks such as ATCC or DMSZ, it is well known that many cell lines have been exchanged between research groups, a situation that increases the probability of CLCC. These problems have already been extensively discussed over the past year, but seem to be ignored by the scientific community. We strongly encourage all scientists to comply with the various recently published guidelines for correct handling of cell lines 37.38

The p53 status in cell lines is now available at the p53 web site (http://p53.free.fr/Database/Cancer_cell_lines/p53_cell_lines.html). A specific section is devoted to cell lines with a controversial p53

2008; Vol. 7 lasue 5

status. We invite all scientists to update these tables with their own findings so that a consensus concerning the p53 status of each cell line can be reached. Finally, we strongly encourage those involved in studies dealing with p53 (or other p53 family members) to regularly check the p53 status of their cell lines.

Material and Methods

Analysis of the biological activity of mutant p53 proteins. Data analysis. The p53 database used for this study contains 21,717 mutations derived from 1,992 publications (UMD p53 database (httpp://p53.free.fr), 2007_R1 release released in January 2007).22 This release contains functional data for the majority of missense p53 mutants. Mutant p53 activity has been described previously.^{22,23} Briefly, 2,314 haploid yeast transformants containing p53 mutations and a GFP-reporter plasmid have been constructed. Mutant p53 activity was tested by measuring the fluorescent intensity of GFP that is controlled by the p21WAF1 promoter sequence of the plasmid after 3 days of growth at 37°C. For functional analysis, frameshift and nonsense mutations were also excluded, as their biological significance has not been clearly established (see text for more information). The mean and 95% Confidence Interval (CI) of the biological activity of all mutants was calculated by using the transactivational activity measured on the p21WAF1 promoter. Similar results were obtained with the activity measured on 7 other promoters of transcription (data not shown).

Statistical analysis. To identify the distinct levels of p53 residual activities among the mutants we used the k-means clustering, ²⁴ whose aim is to partition the data into 3 groups such that the sum of squares from each mutant to the assigned cluster centres is minimized. Three clusters were chosen to represent mutants with no, low and wild-type activity levels. The analysis was based on the measurements of promoter activities of 8 p53 target genes, including p21WAF1, MDM2, BAX, v14-3-3-σ, AIP, GADD45, NOXA and p53R2.

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Note

Supplementary materials can be found at: www.landesbioscience.com/supplement/BerglindCBT7-5-Sup.pdf

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TNF-α-inducing protein, a carcinogenic factor secreted from H. pylori, enters gastric cancer cells

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TNF-\alpha inducing protein (Tip\alpha) is secreted from Helicobacter pylori (H. pylori): it is a potent inducer of TNF-α and chemokine genes, mediated through NF-κB activation, and it also induces tumor-promoting activity in Bhas 42 cells. To investigate the carcinogenic mechanisms of *H. pylori* with Tipα, we first examined how Tipα acts on gastric epithelial cells. We found that fluorescent-Tipα specifically bound to, and then entered, the cells in a dose- and temperature-dependent manner, whereas deletion mutant of Tipa (del-Tipa), an inactive form, neither bound to nor entered the cells, suggesting the presence of a specific binding molecule. Mutagenesis analysis of Tip α revealed that a dimer formation of Tip α with a disulfide bond is required for both specific binding and induction of TNF- α gene expression. A confocal laser scanning microscope revealed some Tip α in the nuclei, but del-Tip α was not present, which indicated that an active form of Tip α can penetrate the nucleus and may be involved in the induction of TNF- α gene expression. Examination of Tipα production and secretion in 28 clinical isolates revealed that *H. pylori* obtained from gastric cancer patients secreted Tipα in significantly higher amounts than did H. pylori from patients with chronic gastritis, suggesting that Tipa is an essential factor in H. pylori inflammation and cancer micro-environment in the human stomach. Tipa is thus a new carcinogenic factor of H. pylori that can enter the nucleus through a specific binding molecule, and its mechanism of action is completely different from that of CagA. © 2008 Wiley-Liss, Inc.

Key words: Τipα; gastric cancer; tumor promoter; NF-κB; TNF-α

Helicobacter pylori has been identified as a causative agent of chronic inflammation, chronic gastritis and peptic ulcer, and is also classified by IARC as a definitive carcinogen for gastric cancer. It is well accepted that persistent H. pylori infection results in an inflammatory response in the stomach by high induction of proinflammatory cytokines, such as tumor necrosis factor-α (TNFα), interleukin-1 (IL-1) and IL-8.2 Based on our previously reported evidence that TNF-α is one of the essential cytokines for tumor promotion and an instigator of a cytokine network sequence of tumor promotion from TNF-α through IL-1 to IL-6 and back to TNF- α , we think a gene product of H. pylori which induces TNF- α plays a significant role in gastric cancer development in humans³⁻⁵: We thus cloned TNF- α inducing protein $(Tip\alpha)$ gene (HP0596) from genome sequence of H. pylori strain 26695. Recombinant Tipα protein strongly induces expression of TNF-α and various chemokine genes, activates NF-κB in mouse gastric epithelial cells MGT-40 and showed in vitro transforming activity of v-H-ras transfected BALB/3T3 (Bhas 42) cells, a standard model of initiated cells. $^{6-9}$ $Tip\alpha$ gene is unique for H. pylori gene nome, which does not have any obvious homologue in other species: only in H. pylori strain was a homologue of Tipa found as H. pylori-membrane protein 1 (HP-MP1) gene from strain SR7791 (94.3% homology) and jph0543 from strain J99 (95.5%). $^{6.10-13}$ Thus, the $Tip\alpha$ gene family codes new carcinogenic factors of H. pylori.

Tipα protein with a molecular weight of 19 kDa is secreted from H. pylori as a homodimer form with 38 kDa. The homodimer form of Tipα is active in the induction of TNF-α gene expression, NF-κB activation in gastric epithelical cells and transforming activity in Bhas 42 cells. Tip α has several unique features: (i) no similarity to other virulence factors of H. pylori, such as vacuolating cytotoxin, immunodominant cytotoxin-associated antigen (CagA) or urease, (ii) secretion from H. pylori in a manner independent of Type IV secretion system, (iii) induction of NF-κB activation in a cag pathogenicity island (cagPAI)-independent manner, (iv) the presence of transforming activity by treatment with a recombinant protein in vitro. Furthermore, we recently found that Tipα has DNA binding activity using surface plasmon resonance assay (Biacore). ¹⁴ All the results were based on the evidence that a recombinant deletion mutant of Tipa (rdel-Tipa), which lacks 6 amino acids including 2 cysteine residues in N-terminal position, was inactive. Therefore, we think that identifying the molecular mechanisms of Tipa will provide a key to understanding the carcinogenic mechanisms of H. pylori infection.

Using fluorescence-labeled rTipa by flow cytometry and immunocytochemical analysis, we first examined how exogenously added recombinant Tipa (rTipa) induces expression of TNF-a and chemokine genes on gastric epithelial cells. Here we report that rTipa bound to cell surface molecules specifically, and then penetrated the cytosol; some rTipa was found localized in the nuclei of gastric cancer cells MGT-40, but rdel-Tipa, an inactive mutant, showed less binding activity and negligible penetration into cytosol and nucleus.

To further investigate the significance of Tipα in gastric cancer development, we next examined Tipa production and secretion in 28 clinical isolates of H. pylori from patients with chronic gastritis and gastric cancer in Saitama Cancer Center, Japan. Although all isolates were positive for Tipa production, we found that H. pylori isolates obtained from gastric cancer patients secreted significantly larger amounts of Tipa into culture broth compared with those from chronic gastritis patients. Therefore, we think that Tipa secreted from H. pylori plays an essential role in inflammation and cancer development in humans.

Material and methods

Preparation of recombinant Tipa and mutant proteins

Recombinant His-tagged proteins of Tipα (rTipα), del-Tipα (rdel-Tipα), cysteine substituted with alanine mutants-Cys5Ala (C5A), Cys7Ala (C7A), Cys5Ala/Cys7Ala (C5AC7A)-were expressed in E. coli transfected pET28(a) + containing each corresponding gene induced with IPTG, and by Ni2+-loaded Hitrap

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chelating column (Amersham Bioscience, Buckinghamshire, UK). ⁶ For generation of cysteine substituted with alanine mutants of Tipα, sense primers were as follows: C5A, 5'-CAGCCA-TATGCTGCAGGCTGCCACTTGCCCAAAC-3'; C7A, 5'-CAG-CCATATGCTGCAGGCTTGCACTGCCCCAAACAC-3'; C5A/C7A, 5'-CAGCCATATGCTGCAGGCTGCCACTGCCCCAAA-CAC-3', and anti-sense primer was 5'-TCTCGGATCCTACAT GGCTATAGGGACTTT-3'. Purities of the 5 recombinant proteins were more than 98% on SDS-PAGE.

Cell culture and reagents

Mouse gastric cancer cell line MGT40 was kindly provided by Dr. Masae Tatematsu, Aichi Cancer Center Research Institute, Aichi, Japan. MGT-40 cells were maintained in DMEM with 10% fetal bovine serum (JRH Biosciences, KS) and MITO+ serum extender (Becton Dickinson Labware, MA), as described previously.¹⁵ Anti-Tipα antibody was obtained by immunization of rabbit with 19-mer synthetic peptide, also as described previously.⁶ Anti-HSP90, anti-EGF receptor and anti-lamin B antibodies were purchased from Santa Cruz (CA).

H. pylori strains and culture conditions

Twenty-eight *H. pylori* clinical isolates were obtained from patients with chronic gastritis (11 patients) and gastric cancer (17 patients), using M-BHM *H. pylori* selection agar plates (Nissui Pharmaceutical Co., Tokyo, Japan) at Saitama Cancer Center between 2001 and 2006. *H. pylori* strain 26695 were kindly provided by Dr. Chihiro Sasakawa (The Institute of Medical Science University of Tokyo, Tokyo Japan), and ATCC43504 was purchased from American Type Culture Collection (VA). All *H. pylori* isolates were cultured in Brucella broth (Becton Dickinson Microbiology Systems, MD) containing 10% horse serum (Nippon Bio-Test Lab, Tokyo, Japan) at 37°C, with shaking in microaerobic conditions, as described previously. ^{6,16}

Western blotting

After 3 days culture of H. pylori strains, culture broths with volumes (7.5–15 μ l) equal to 1.0 OD₅₄₀ of bacteria amount were separated on 12% SDS-polyacrylamide gel in the presence of dithiothreitol (DTT), and blotted onto nitrocellulose membranes. Tip α protein was visualized by the ECL detection system (Amersham Bioscience) using anti-Tip α antibody. Amounts of Tip α were measured by density of a band of 19 kDa (Tip α) by NIH Image, and expressed as a relative unit based on Tip α amount in culture broth of 26695 strain. The results were expressed as the average of 2 separate experiments.

Expression of TNF-α gene

MGT-40 cells were treated with recombinant proteins for 1 hr, and total RNA was isolated with ISOGEN reagent (Nippon Gene, Tokyo, Japan). Expression of *TNF*-α gene was determined by semiquantitative RT-PCR, as described previously.

Flow cytometry

rTipα was labeled with FITC using EZ-labelTM FITC protein labeling kit (Pierce Biotechnology, IL) according to the manufacturer's instructions. MGT-40 cells (1 \times 10⁶ cells/ml) in PBS were incubated with various concentrations of FITC-rTipα at 4°C for 2 hr. Cells were then washed with PBS and analyzed by flow cytometry (EpicsXL, Beckman Coulter, Tokyo, Japan). FITC-labeled BSA was used as negative control. Equal absorbance of FITC-rTipα or FITC-BSA at 495 nm (1 mol FITC/ mol protein) was used in the experiments. Binding of FITC-rTipα to MGT-40 cells was measured as mean fluorescence intensity (FI). ¹⁷ Binding of FITC-rTipα (2.5 μM) to MGT-40 cells was also analyzed in the presence of nonlabeled rTipα, rdel-Tipα, C5A, C7A or C5AC7A mutant at various concentrations.

Cytochemical analysis

rTipα, rdel-Tipα and BSA were labeled with Alexa Fluor 488 protein labeling kit (Invitrogen, Tokyo, Japan) according to the manufacturer's instructions. MGT-40 cells were incubated with Alexa Fluor 488-labeled rTipα (AF488-Tipα), rdel-Tipα (AF488-del-Tipα) or BSA (AF488-BSA) at a concentration of 5.0 μM, with equal absorbance at 495 nm for 1 hr. After fixation with 4% paraformaldehyde containing 0.2% Triton X-100, the cells were observed using fluorescence microscope (Leica Microbiosystems, Tokyo, Japan).

Immunocytochemical analysis

MGT-40 cells were incubated with rTipα (100 μg/ml) for 1 hr at 37°C, and then washed with PBS. The cells were fixed with 4% paraformaldehyde containing 0.2% Triton X-100. After blocking with Block Ace (Dainippon Pharm., Osaka, Japan), the cells were treated with anti-Tipα antibody, and stained with Alexa Fluor 488-conjugated anti-rabbit IgG (Invitrogen, Tokyo, Japan). Then the cells were stained with propidium iodide and analyzed using confocal laser scanning microscope LSM 5 PASCAL (Carl Zeiss, Germany).

Subcellular fraction analysis

MGT-40 cells were incubated with rTip α or rdel-Tip α (5 μ M) for 1 hr at 37°C and washed with PBS, and the cells were fractionated into cytosol, membrane and nuclei using Qproteome cell compartment kit (Qiagen, Düesseldorf, Germany) according to the manufacturer's instructions. ¹⁸ Seven micrograms of each fraction was subjected to Western blotting, and analyzed to detect rTip α and rdel-Tip α proteins by anti-Tip α antibody, and heat shock protein 90 (HSP90; a marker for cytosol), epidermal growth factor receptor (EGFR; for membrane) and lamin B (for nucleus) as the controls for cell fractionation by corresponding antibodies.

Statistical analysis

The differences in secreted amounts of $\text{Tip}\alpha$ from H. pylori obtained from patients with gastritis and gastric cancer were analyzed by Mann-Whitney analysis, with exact p value using SPSS V14.0 (SPSS, Chicago, IL).

Results

Specific binding of FITC-labeled Tipa protein to MGT-40 cells

We first examined specific binding of FITC-labeled rTipα to MGT-40 cells using flow cytometry: Incubation of MGT-40 cells with FITC-rTipa protein at 4°C for 2 hr significantly increased cellular fluorescence dose-dependently (Fig. 1a). Quantitative analysis of "mean cellular FI") indicated that binding of FITCrTipα to MGT-40 cells was significant and saturated at concentrations of 5.0-7.5 µM. However, FI in the case of FITC-BSA was significantly low and not saturated up to 7.5 µM (Fig. 1a). Furthermore, the binding of FITC-rTipa to MGT-40 cells was dosedependently inhibited with nonlabeled rTipa, as shown in Figure 1b. However, rdel-Tipα, an inactive protein lacking 6 amino acids, including 2 cysteine residues, showed less inhibitory activity. Concentration of 50% inhibition (IC50) values are 1.9 µM for rTipα and 20.0 μM for rdel-Tipα. The inhibition of specific binding with nonlabeled rTipa was about 10 times stronger than with rdel-Tipa. These results correlated closely with induction of TNFα gene, as well as biological activities such as induction of NF-κB activation and chemokine gene expressions in MGT-40 cells, and in vitro transformation of Bhas 42 cells, as reported previously.

To clarify the significance of cysteine residues more precisely, we made 3 mutants: One cysteine substituted with an alanine at 5 or 7 position (C5A or C7A mutant), and two cysteines substituted with 2 alanines at 5 and 7 positions at the same time (C5A/C7A double mutant) (Fig. 2a). Each recombinant protein was analyzed by SDS-PAGE in the absence of DTT. C5A and C7A proteins formed a dimer similar to Tipα, since 1 cysteine residue still

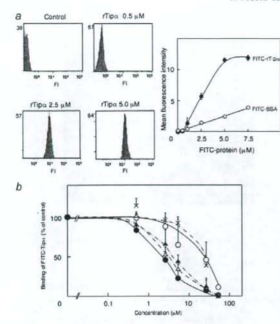


FIGURE 1 – Binding of FTTC-rTip α to MGT-40 cells analyzed by flow cytometry. (a) MGT-40 cells were incubated with various concentrations of FTTC-rTip α for 2 hr at 4°C. Fluorescence intensity (Ft) was detected by Flow cytometer, as described in Material and Methods. FTTC-BSA was used as a control. (b) MGT-40 cells were incubated with FTTC-rTip α (2.5 μ M) in the presence of nonlabeled rTip α (\bullet), C5A (\triangle), C7A (\bullet), C5A/C7A (X) double mutant or rdel-Tip α (\bigcirc). Fluorescence intensity in the absence of nonlabeled protein is expressed as 100%. The results are the average of 2 independent experiments. Bars show SD.

remained, and C5A/C7A double mutant protein showed only a monomer band with a molecular weight of 21 kDa, with 2 cysteines entirely replaced (Fig. 2b). C5A and C7A mutant proteins dose-dependently inhibited the binding of FITC-rTip α to MGT-40 cells similar to rTip α , and C5A/C7A double-mutant proteins showed weak inhibitor activity (Fig. 1b). IC $_{50}$ values were 2.4 μ M for C5A, 2.9 μ M for C7A and 21.0 μ M for C5A/C7A. Their binding activities to MGT-40 cells were identical: treatment with C5A and C7A proteins strongly induced TNF- α gene expression, but C5A/C7A protein did not. These results indicated that homodimer formation with a disulfide bond through a cysteine residue is essential for Tip α binding to MGT-40 cells as well as TNF- α induction and carcinogenic activities (Fig. 2c).

The results suggest the presence of a specific binding molecule which can recognize the homodimer form of Tipo on the cell surface of MGT-40 cells. Specific binding of FITC-rTip α was also confirmed in human gastric cancer cell lines (MKN-1, MKN-45 and MKN-74) by flow cytometry (data not shown), and determination of the specific binding molecule(s) is now under investigation. When incubation of the cells with FITC-rTip α was conducted at 37°C, binding of Tip α was lower than that incubated at 4°C, and not saturated, although bindings of FITC-BSA were the same in both cases. The results led us to believe that rTip α internalizes into MGT-40 cells in a temperature-dependent manner.

Incorporation of Tipa into MGT-40 cells

As the fluorescence of FTTC is quite unstable for fluorescence microscope analysis, rTip α was labeled with a photostable dye, Alexa Fluor 488. After incubation of MGT-40 cells with Alexa

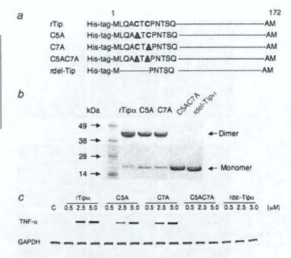


FIGURE 2 – Significance of a cysteine residue for homodimer formation and TNF- α gene induction. (a) Schematic representation of Tip α , cysteine-substituted-mutants and rdel-Tip α . (b) SDS-PADE analysis of recombinant proteins in the absence of DTT. rTip α . C5A and C7A formed homodimers, but C5A/C7A double mutant and rdel-Tip α did not. (c) Induction of TNF- α gene expression in MGT-40 cells. Expression of TNF- α gene was examined by semiquantitative RT-PCR, as described in Material and Methods.

Fluor 488-conjugated Tipα (AF488-Tipα) at 37°C, significant fluorescence was observed in the cells after 10-min incubation, it increased time-dependently until 30 min, and was sustained until 2 hr. Figure 3a shows strong fluorescent spots in MGT-40 cells after 1-hr incubation with AF488-rTipα at 37°C. However, the cells incubated with AF488-rdel-Tipα showed only slight fluorescence similar to those incubated with AF488-BSA, although the same "fluorescence intensities" of AF488-labeled proteins were used in these experiments. Fluorescence in the cells incubated with AF488-Tipα increased in dose-dependent and time-dependent manners. All the results show that rTipα significantly incorporated into the cells, while the inactive mutant protein rdel-Tipα did not.

To confirm incorporation of rTip α protein into the cells, we next conducted immunocytochemical analysis with anti-Tip α antibody, and observed the results under a confocal laser scanning microscope. Anti-Tip α antibody clearly recognized Tip α protein in the cytosol of the cells after treatment with Tip α for 1 hr (Fig. 3b), but nonimmunized rabbit IgG did not show any significant fluorescence in rTip α -treated cells. Nontreated cells were negatively stained. All results indicate that Tip α enters the gastric epithelial cells, resulting in expression of TNF- α and Chemokine genes.

Localization of rTipa in nucleus

Confocal laser scanning microscope analysis revealed that some fluorescent spots that had reacted with fluorescent anti-Tipα antibody were located in the nuclei of MGT-40 cells treated with Tipα, revealing the internalization of Tipα into nuclei (Fig. 4a). To confirm localization of rTipα in nuclei, we next conducted cell fractionation into cytosol, membrane and nuclei, after treatment with rTipα for 1 hr at 37°C. As shown in Figure 4b, rTipα was present in each cell fraction compared with marker proteins of each separated fraction: HSP90 for cytosol, EGFR for membrane and lamin B for nucleus. Based on the density of protein bands, we calculated that about 10% of Tipα had localized in the nuclei of the cells. However, while small amounts of rdel-Tipα, an inactive mutant, were detected in cytosol and membrane fractions—

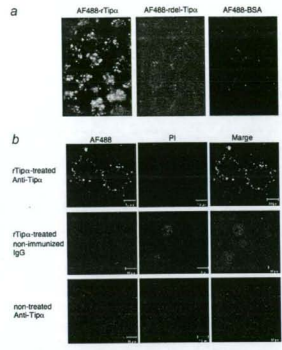
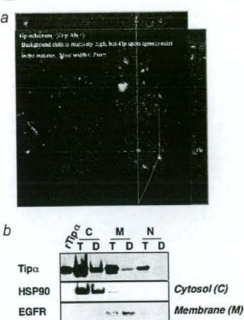


FIGURE 3 – Incorporation of Alexa Fluoro 488 (AF488)-labeledrTipα into MGT-40 cells. (a) After incubation with AF488-rTipα, AF488-rdel-Tipα or AF488-BSA at a concentration of 5.0 μM for 1 hr at 37°C, MGT-40 cells were washed with PBS and then observed using fluorescence micoroscope. (b) Existence of rTipα in the cytosol was confirmed with confocal laser scanning microscope after immunocytochemical staining. MGT-40 cells were incubated with rTipα (5.0 μM) for 1 hr at 37°C, then subjected to anti-Tipα antibody or nonimmunized rabbit IgG (for a negative control) followed by AF488-labeled anti-rabbit IgG, as described in Material and Methods. As control, nontreated cells were subjected to anti-Tipα antibody. Nuclei were stained with propidium iodide (PI).

about 13.5–17.5% of rTip α —none in the nuclei fraction was found. These results strongly supported the results by confocal laser scanning microscope. Recently, we found that Tip α directly bound to DNA, but del-Tip α did not. We therefore think that Tip α in the nuclei plays an essential role in the induction of TNF- α and chemokine genes and carcinogenic activity. This is the first report that a protein secreted from H. pylori enters the nuclei, and these results will provide a deep insight in understanding the mechanisms of gastric cancer development by H. pylori infection.

Large amounts of Tipa protein secreted from H. pylori from gastric cancer patients

To further understand the significance of Tip α in human gastric cancer development, we examined differences of production and secretion of Tip α among various H. pylori clinical isolates from biopsy samples of gastric mucosa obtained from patients with chronic gastritis and gastric cancer. Twenty-eight H. pylori isolates, from 17 gastric cancer patients and 11 chronic gastritis patients, were cultured in Brucella, broth containing 10% horse serum at 37°C in microaerobic condition for 3 days after inoculation of 5×10^8 CFU. All isolates produced CagA, which coincided with previous results in which H. pylori isolated from Japanese patients were more than 98%-positive cagPAI. 19 All H. pylori isolates produced Tip α protein, and Tip α in bacterial extracts was consistently present (data not shown). In contrast, the amounts of



Lamin B

82

FIGURE 4 – Nuclear localization of rTipα. (a) Some yellow fluorescent spots were observed in the nuclei using confocal laser scanning microscope. (b) After cellular fractionation of rTipα or rdel-Tipα-treated cells into the cytosol, membrane and nuclei, and each fraction was analyzed by Western blotting with anti-Tipα antibody. Each fraction was confirmed by detection of specific marker proteins—HSP90 for cytosol, EGFR for membrane and lamin B for nuclei—by anti-HSP90, anti-EGFR and anti-lamin B antibodies, respectively. The gel stained with quick CBB is also indicated.

Nuclear (N)

Tip α in culture broth varied, and clinical isolates obtained from gastric cancer patients secreted Tip α protein significantly more than did those from gastritis patients (Figs. 5a and 5b).

H. pylori 26695 strain, from which genome $Tip\alpha$ gene was cloned, secreted about 1.0 ng of $Tip\alpha/10^\circ$ CFU/ml, and thus was expressed as 1 relative unit. Clinical isolates from cancer patients secreted $Tip\alpha$ protein at 1.4–13.4 relative units, and those from gastritis patients secreted at 0.8–6.7 relative units (Fig. 5b). Interestingly, H. pylori isolated from 3 of 11 gastritis patients who developed gastric cancer also secreted larger amounts of $Tip\alpha$, similar to those from cancer patients. Their median values are 2.0 for H. pylori from 8 patients with gastritis, 3.8 from 3 patients who later developed cancer and 4.5 from 17 patients with gastric cancer. The difference of secreted amounts of $Tip\alpha$ between 17 gastric cancer patients and 8 chronic gastritis patients (excluding those that developed cancer) is statistically significant (p = 0.004), although the difference was not so great. These results

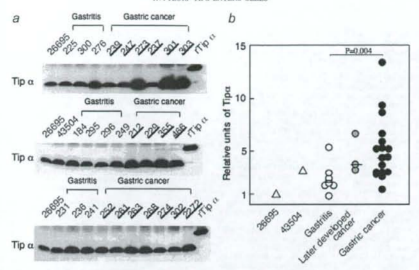


FIGURE 5 — Large amounts of Tip α were secreted from H. pylori clinical isolates obtained from gastric cancer patients. (a) Eleven clinical isolates from chronic gastritis patients, including 3 patients, who later developed gastric cancer, and 17 isolates from cancer patients were cultured for 3 days in microaerobic conditions, after inoculation of 0.5 McFalrand in 8-ml Brucella broth with 10% horse serum. Aliquots of culture broth equivalent to 1.0 OD_{540} were analyzed by Western blotting with anti-Tip α antibody, as described in Material and Methods. (b) Relative units of Tip α in culture broth were measured for intensity of bands by NIH image, with Tip α in culture broth of 26695 strain used as a control, and expressed as 1 relative unit. Each dot corresponds to Tip α in culture broth from each clinical isolate (α) from chronic gastritis patients, gray circle from patients later developed gastric cancer, from gastric cancer patients). Bars indicate the median value of each group. Difference between gastritis patients (excluding those that developed cancer) and gastric cancer patients was statistically significant (α) = 0.004).

indicate that secreted $\text{Tip}\alpha$ plays a significant role in gastric cancer micorenvironment during H, pylori infection in humans.

Discussion

This manuscript demonstrated that Tip α protein is a new protein secreted from H. pylori: It can enter the gastric epithelial cells, where it then localizes in the nuclei of the cells. As for the specific function of Tip α protein in the nucleus, we recently reported that Tip α protein has direct DNA binding activity as determined by surface plasmon resonance (Biacore) assay, and that a homodimer of Tip α bound to DNA oligomer more strongly than did a monomer of del-Tip α . Homodimer formation of Tip α is thus necessary for penetration into nucleus and also for the DNA binding activity. Interestingly, a dimer of Tip α dose-dependently bound to 9 bases of DNA oligomers in TNF- α promoter sequence. Pretreatment with N-acetylcysteine significantly inhibited penetration of Tip α into MGT-40 cells and induction of TNF- α gene expression (unpublished results). Therefore, we think that Tip α protein in nucleus regulates expression of TNF- α and IL-6 genes as well as chemokine genes, and thus plays a crucial role in carcinogenic activity.

How Tipα protein secreted from *H. pylori* enters the gastric epithelial cells remains to be clarified. We think that receptor-mediated endocytosis is involved, based on results showing that homodimers of Tipα, C5A and C7A proteins specifically bound to the MGT-40 cells more strongly than monomers of del-Tipα and C5A/C7A proteins did. Recently it was reported that decay-accelerating factor (DAF), a glycoprotein, might act as a receptor for *H. pylori*: a mediator of gastric inflammation and genetic deficiency of DAF attenuated the development of inflammation among *H. pylori*-infected mice.²⁰

After Tipα binds to the protein, Tipα trafficking may occur by caveolae or by lipid raft. It has been shown that some proteins—termed translocatory proteins—such as human immunodeficiency virus Tat, can enter the cell by endocytosis into endocytic vesicles and translocate into the nuclei, but the mechanisms are not yet clear. ^{18,21} We believe that determining the mechanisms of Tipα trafficking will provide new insight into carcinogenic mechanisms with *H. pylori* infection and various virulence factors.

Our findings showing that $H.\ pylori$ obtained from gastric cancer patients secreted Tip α protein significantly more than those from chronic gastritis patients did indicate that larger amounts of Tip α protein induce stronger expression of $TNF-\alpha$ and chemokine gene expression in the gastric mucosa. Recently, Inoue et al. reported nasal vaccination with Tip α significantly reduced the inflammatory cytokines TNF- α and IL-12 in gastric mucosa of mice infected with $H.\ pylori.^{22}$ Cao et al., using Mongolian gerbils, reported a severity of chronic gastritis associated with higher levels of mRNA of TNF- α and IL-1 β , with pathogen $H.\ pylori$ linked to glandular gastric carcinogenesis. Thus, Tip α may induce cancer development through a unique mechanism that is completely different from that of CagA with Type IV secretion system.

All the results show that $Tip\alpha$ is a unique carcinogenic factor of H. pylori and is both a suitable marker for detection of high risk in gastric cancer and a molecular target for cancer prevention.

Acknowledgements

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Microsatellite instability-low colorectal cancer acquires a KRAS mutation during the progression from Dukes' A to Dukes' B

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The classification of colorectal cancer (CRC) by microsatellite instability (MSI) status is important for effective clinical management. In fact, microsatellite instability-high (MSI-H) cancer has distinctive clinicopathological and molecular features. However, microsatellite instability-low (MSI-L) cancer is not clearly defined. The objective of this study was to further clarify the characteristics of MSI-L CRC. A consecutive series of 940 primary CRCs were subdivided into three groups according to the level of MSI and analyzed the clinicopathological features and genetic changes in the KRAS, BRAF and p53 mutation and the loss of heterozygosity (LOH) of adenomatous polyposis coli (APC) gene and methylation status of the O6-methylguanine-DNA methyl-25 transferase (MGMT) and MLH1 promoter. Of the 940 CRCs, 5.9% were MSI-H, 7.1% were MSI-L and 87% were microsatellite stable (MSS), KRAS and BRAF mutations were detected in 39.4 and 4.6% of the CRCs, respectively. The frequency of KRAS mutations in MSI-H, MSI-L and MSS cancer was 30, 48 and 39 %, 30 respectively. The proportion of KRAS mutations in MSI-L cancer increased from 16 to 63% accompanying the progression from Dukes' A to Dukes' B. While the LOH of D5S346, which is located near the APC gene, and p53 mutation was observed in 75 and 67% of MSI-L CRC at Dukes' A, respectively. These results indicated that the LOH of APC and p53 mutation has already occurred by the Dukes' A lake 'suppressor pathway' but not the KRAS mutation in MSI-L CRCs. The genes involving MSI-L carcinogenesis are similar to MSS but the timing and frequency of the KRAS mutation is different.

an Introduction

There are two types of genomic instability, microsatellite instability (MSI or MIN) or chromosomal instability associated with the carcinogenesis process of colorectal cancer (CRC) (1). The great majority of CRCs develop through the chromosomal instability pathway (also called the 'suppressor pathway'), which arise from adenomas and is initiated by the inactivated adenomatous polyposis coli (APC) gene and followed by the well-established genetic steps involved in the adenoma-carcinoma sequence (2,3). While another type of genomic instability, MSI caused by a failure of the DNA mismatch repair (MMR) system, is observed in ~10% of all CRCs (4-7). DNA MMR deficiency leads mutations in the target genes that are implicated in tumor progression such as TGFbetaR2 (8), IGF2R (9), CDX2 (10) and BAX (11) and it is known as the 'mutator pathway'. MSI can be subdivided into three groups, microsatellite instability-high (MSI-

Abbreviations: APC, adenomatous polyposis coli; CI, confidence interval; CRC, colorectal cancer; HR, hazard ratio; LOH, loss of heterozygosity; MGMT, O*-methylguanine-DNA methyltransferase; MMR, mismatch repair; MSI, microsatellite instability; MSI-H, microsatellite instability-high; MSI-L, microsatellite instability-low; MSS, microsatellite stable; PCR, polymerase chain reaction.

H), microsatellite instability-low (MSI-L) and microsatellite stable 55 (MSS), according to the degree of instability. The recommended method to distinguish these subgroups is to analyze paired tumor and normal tissue DNAs using a panel of five microsatellite markers known as the Bethesda panel (12).

The MSI-H CRCs phenotype is more likely to occur at a proximal site, to occur in women, to be associated with a favorable prognosis (5,7,13–15) and severe inflammatory cell infiltration into the tumor tissue (16,17). A large percent of MSI-H CRC is sporadic and demonstrates somatic promoter methylation of the hMLH1 gene (18,19), whereas a germ line mutation of the MMR genes, such as hMSH2, 65 hMLH1, hMSH6 and hPMS2, is found in the majority of MSI-H CRC without hMLH1 promoter methylation and is known as Lynch syndrome/hereditary non-polyposis colorectal cancer (20–23). Recent morphological and molecular studies have proposed the existence of a serrated pathway, thus suggesting that serrated polyps may serve as a precursor of the MSI-positive cancers (24–26).

On the other hand, most studies have found no obvious clinicopathological or molecular differences between MSI-L and MSS cancers (27). The DNA MMR genes hMLH1 and hMLH2 do not appear to be implicated in the MSI-L subset (28). Some studies have reported that MSI-L is associated with cancers from individuals with germ line mutations of hMLH6 (29), but genetic alteration of this gene is infrequent in MSI-L CRC patients. Furthermore, some researchers deny the presence of MSI-L cancers because most non-MSI-H cancers exhibit MSI-L when large numbers of microsatellite loci are tested (27,30).

Meanwhile, there is evidence indicating that the MSI-L phenotype could reflect a distinct pathway of tumor development with a different clinical behavior and different genetic and epigenetic changes. For example, a high frequency of a KRAS mutation (31,32) that is associated with loss of expression of the O°-methylguanine-DNA methyltransferase (MGMT) gene by methylation of its promoter region (33), lower frequency of 5qLOH (31), a high frequency of APC mutation (34) and reduced expression of Bel-2 protein (35) are global molecular phenotypes by which MSI-L cancers are distinguished from non-MSI-L cancers.

Therefore, MSI-L CRC is still controversial. This study investigated the genetic changes and clinicopathological features of MSI-L CRCs using a series of 940 CRCs.

Materials and methods

Patients and tissue samples

A consecutive series of 940 primary CRCs excised surgically at Saitama Cancer Center from January 1998 to May 2006 were investigated after obtaining the informed consent. Any patients who were treated by preoperative radiotherapy or chemotherapy were excluded. Furthermore, patients with inflammatory bowel disease or a known history of familial adenomatous polyposis were also excluded. This study was approved by the Ethics Committee of the Saitama Cancer Center.

Analysis of MSI

Primary CRCs and paired normal colorectal mucosa obtained by surgery were immediately frozen at -80°C . The genomic DNA was extracted from fresh frozen specimens using standard methods. The Bethesda five markers, BAT25, BAT26, D5S346, D25123 and D175250, were used to classify the MSI status of the tumors. Polymerase chain reaction (PCR) and subsequent analyses were performed as reported previously (5), CRCs were subdivided into three groups according to the degree of MSI; MSI-H if two or more of the five markers show instability, MSI-L if only one marker shows instability and MSS if absence of MSI in all five markers. MSI-positive markers were re-examined at least twice to confirm the result. Loss of heterozygosity (LOH) was defined by at least a 30% reduction in the relative intensity of one allele in the tumor in comparison with normal levels.

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Analysis of KRAS, BRAF and p53 mutation

The mutations in exon 1 and 2 of the KRAS gene were analyzed by denaturing gradient gel electrophoresis as described previously (36).

The BRAF V600E mutation was examined using PCR combined with restriction enzyme digestion. DNA fragments containing exon 15 of the BRAF gene were amplified by PCR using the following oligonucleotide primers: BRAF forward primer 5'-CTGTTTCATACTTACTACACC-3' and BRAF reverse primer 5'-CTGTTCAAACTGATGGGACC-3'. PCR amplification was carried out with 100 ng of genomic DNA in a volume of 20 μl containing 0.2 μM deoxynucleoside triphosphate, 0.1 μM each of primers and 1 U Taq Gold. Thermal cycling was initiated with denaturation at 94°C for 10 min followed by 37 three-step cycles at 94°C for 30 s, 59°C for 30 s and 72°C for 45 s and followed by a final incubation for 7 min at 72°C. PCR products were digested with HpyCH4III at 37°C for 1 h and analyzed on 8% polyacrylamide gels.

The mutations in exon 5-8 of p53 gene were analyzed by denaturing gradient gel electrophoresis as described previously (37).

Analysis of hMLH1 and MGMT promoter methylation

hMLH1 and MGMT promoter methylation was analyzed in 55 of MSI-H cancer and 67 of MSI-L cancer samples. The methylation status of each gene was determined by the methods previously reported (5). The primers were hMLH1 methylation specific, 5'-AACGAATTAATAGGAAGAGGCGGATAGCG-3' and 5'-CGTCCCTCCCTAAAACGACTACTACCC-3'; hMLH1 unmethylation specific, 5'-TAAAAATGAATTAATAGGAAGAGTGGATAGTG-3' and 5'-ATCTCTTCATCCCTCCCTAAAACA-3'; MGMT methylation specific, 5'-TTTCGACGTTCCTCCCTAAAACA-3' and 5'-GCACTCTTCCGAAAA-

CGAAACG-3' and MGMT unmethylation specific, 5'-TTTGTGTTTTGA-TGTTTGTAGGTTTTTGT-3' and 5'-AACTCCACACTCTTCCAAAAACA AAACA-3'.

143

Statistical analysis

Differences were assessed using the chi-square or Fisher's exact test for categorical variable and unpaired Student's *t*-test for continuous factors. The overall survival was defined as the interval from the date of resection until the date of death from any cause, censored patients being those alive at the close of the study or lost to follow-up. Survival was measured from the date of the resection of the CRCs until death or until the censor date of 1 July 2006. The distribution of survival time was compared with the use of the log-rank test; survival distribution curves were estimated by the method of Kaplan-Meier. Multivariate analyses were performed with the use of the Cox proportion hazard model. The independent prognostic factors for survival were determined by a stepwise backward conditional selection in which the non-significant factors (P > 0.1) were successively rejected. All statistical analyses were performed using the StatView 5.5 program. P < 0.05 was considered to be statistically significant in all cases.

Results 160

MSI status

The Bethesda panel, BAT25, BAT26, D5S346, D2S123 and D17S250, was used to classify the MSI status of the tumors. Of the 940 CRCs, 55 (5.9%) were MSI-H, 67 (7.1%) were MSI-L and 818 (87%) were MSS (Table 1). Mononucleotide marker BAT25 and BAT26 exhibited

Table I. Clinicopathological and genetic features of CRCs

	MSS, n (%)	SS, n (%) MSI-L, n (%)	MSI-H, n (%)	P value		
				MSI-L versus MSS	MSI-H versus MSS	MSI-L versus MSI-H
Patient	818 (87.0)	67 (7.1)	55 (5.9)	0.3723	0.0011	0.0661
Men	509 (62.2)	38 (56.7)	22 (40)			
Women	309 (37.8)	29 (43.3)	33 (60)			
Mean (±SE) age	63.6 ± 10.3	63.3 ± 9.95	60.5 ± 13.4	0.785	0.035	0.199
Location				0.44	< 0.0001	0.0003
Proximal	217 (26.5)	22 (32.8)	36 (65.5)			
Distal	243 (29.7)	16 (23.9)	12 (21.8)			
Rectum	358 (43.8)	29 (43.3)	7 (12.7)			
Tumor size				0.23	< 0.0001	< 0.0001
Mean ± standard error (mm)	45.42 ± 24.1	41.8 ± 23.0	61.35 ± 32.3			
Histologic feature				>0.999*	< 0.0001*	0.0018 ^a
Well differentiated	73 (8.9)	10 (14.9)	4 (7.3)			
Moderately differentiated	706 (86.3)	54 (80.6)	38 (69.1)			
Poorly differentiated	16 (2.0)	3 (4.5)	10 (18.2)			
Mucinous	21 (2.6)	0 (0)	2 (3.6)			
Others	2 (0.2)	0 (0)	1(1.8)			
Mucinous component		A CONTRACTOR		0.73	< 0.0001	0.001
+	84 (10.3)	6 (9.0)	18 (32.7)			
-	734 (89.7)	61 (91.0)	37 (67.3)			
Dukes' stage				0.22	0.012	0.039
A	150 (18.3)	19 (28.4)	12 (21.8)			
В	246 (30.1)	16 (23.9)	26 (47.3)			
C	246 (30.1)	20 (29.9)	13 (23.6)			
D	176 (21.5)	12 (17.9)	4 (7.3)			
Depth of tumor invasion	CHARLESONE	400000000000000000000000000000000000000	14558	0.27	0.073	0.42
TI	68 (8.3)	8 (11.9)	5 (9.1)	1,000.00	10000	
T2	123 (15.0)	14 (20.9)	7 (12.7)			
T3	571 (69.8)	39 (58.2)	34 (61.8)			
T4	56 (6.8)	6 (9.0)	9 (16.4)			
Extramural venous invasion			4.45	0.99	0.69	0.76
+	586 (71.6)	48 (71.6)	38 (69.1)	139550	235	77.00
_	232 (28.4)	19 (28.4)	17 (30.9)			
KRAS mutation	100.17			0.1836	0.1798	0.051
+	311 (39.5)	32 (47.8)	16 (30.2)	100 to 10		
T	477 (60.5)	35 (52.2)	37 (69.8)			
BRAF mutation	(0010)	()	(00-10)	0.3885	< 0.0001	< 0.0001
+	21 (2.7)	3 (4.5)	17 (32.1)	0.000	- United I	10.0001
1	767 (97.3)	64 (95.5)	36 (67.9)			

^aWell and moderately differentiated versus mucinous and poorly differentiated.