Satellite RNAs promote pancreatic oncogenic processes via the dysfunction of YBX1

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Highly repetitive tandem arrays at the centromeric and pericentromeric regions in chromosomes, previously considered silent, are actively transcribed, particularly in cancer. This aberrant expression occurs even in K-ras-mutated pancreatic intraepithelial neoplasia (PanIN) tissues, which are precancerous lesions. To examine the biological roles of the satellite RNAs in carcinogenesis, we construct mouse PanIN-derived cells expressing major satellite (MajSAT) RNA and show increased malignant properties. We find an increase in frequency of chromosomal instability and point mutations in both genomic and mitochondrial DNA. We identify Y-box binding protein 1 (YBX1) as a protein that binds to MajSAT RNA. MajSAT RNA inhibits the nuclear translocation of YBX1 under stress conditions, thus reducing its DNA-damage repair function. The forced expression of YBX1 significantly decreases the aberrant phenotypes. These findings indicate that during the early stage of cancer development, satellite transcripts may act as ‘intrinsic mutagens’ by inducing YBX1 dysfunction, which may be crucial in oncogenic processes.
Pancreatic cancer, one of the most intractable diseases, develops in incremental steps with the sequential activation of oncogenes and the dysfunction of tumour suppressor genes\(^2\,3\). However, the frequently mutated genes are relatively limited, such as KRAS, TP53, CDKN2A, SMAD4 (refs 3–7). In particular, constitutively active mutations of the K-ras gene are observed in almost all pancreatic cancers (>95%) and are found in 36–87% of pancreatic intraepithelial neoplasia (PanIN) tissues, which are considered to be the precancerous lesions of the pancreatic cancer\(^8\,9\). These observations might indicate that mutations in K-ras occur during the earlier stage of pancreatic PanIN-carcinoma sequence, and the accumulation of mutations in other genes during the later stage causes the cellular transformation. These hypotheses are supported by the fact that genetically engineered mice with pancreas-specific K-ras mutation form regional PanIN-like lesions via acinar-to-ductal metaplasia, whereas the additional deletion of tumour suppressor genes, such as TP53, SMAD4 or TGFβ2, causes the development of invasive cancer\(^9\).

Satellite DNAs, which consist of highly repetitive non-coding sequences in huge monomeric arrays, are largely located in the centromeric and pericentromeric regions of the chromosomes. These chromosomal structures are conserved in almost all eukaryotes, although each monomeric sequence is different between species\(^10\). In the mouse genome, the centromeric region consists of 120-base monomeric arrays, called minor satellites, and the pericentromeric region is composed of 234-base monomeric arrays, called ‘major satellites’ (MajSATS)\(^9\). Previously, satellite regions were believed to be silent because of their constitutive heterochromatin structures. However, recent studies have provided evidences that these regions are actively transcribed\(^11\). Some reports have shown that the appropriate transcription of these satellite regions is essential for accurate cell division\(^12\,13\), heterochromatin establishment in mouse embryonic development\(^14\,15\) and cell differentiation\(^16\). In contrast to these physiological roles, the aberrant transcription of satellite sequences can be observed in epithelial tumours, especially in pancreatic cancers including PanIN lesions\(^18\). While the overexpression of satellite RNAs may cause mitotic errors, such as centrosome amplification and incorrect separation, or genomic DNA damage, such as double-strand breaks\(^15\,19\), the pathological roles of these aberrantly expressed satellite RNAs, especially in precancerous tissues, are not yet fully determined.

Y-box binding protein 1 (YBX1) is a multifunctional protein, mainly known as a transcriptional and translational regulator that is involved in DNA repair, centrosome maturation and mRNA splicing\(^21\,22\). This protein is typically localized to the cytoplasm and acts as an RNA-binding protein\(^23\). However, when cells are exposed to stress conditions, such as oxidative stress and ultraviolet irradiation, YBX1 often translocates into the nucleus\(^24\,25\). Nuclear YBX1 has been considered to participate in DNA-damage repair activity via diverse but currently undefined mechanisms\(^22\).

In this study, we confirmed that MajSAT RNA is expressed in precancerous PanIN lesions in vivo. This finding led us to hypothesize that the aberrant expression of MajSAT RNA contributes molecularly to malignant transformation. We show that aberrantly expressed MajSAT RNA in PanIN-derived cells increases the chromosomal instability and mutations in genomic and mitochondrial DNAs (mtDNA), by inducing YBX1 dysfunction. The increased instability and mutations due to satellite RNAs may molecularly explain the process of the transformation of precancerous cells to invasive cancers.

**Results**

**Expression of MajSAT RNA in mouse pancreatic cells.** First, we examined the MajSAT RNA expression status in two types of genetically induced mouse pancreatic tumour models. Mice with constitutively active K-ras specifically in the pancreatic cells (KrasG12D mice) spontaneously develop PanIN lesions resembling human PanIN tissues\(^2\). Mice with constitutively active K-ras and Tgfβ2r deletion specifically in the pancreatic cells (KrasG12D + Tgfβ2r\(^−/−\) mice) develop aggressive pancreatic carcinoma at 6–7 weeks of age\(^2\). Consistently with the previous report\(^18\), MajSAT RNA was expressed in cancer cells as well as in PanIN cells, while no MajSAT RNA expression was observed in wildtype tissues (Fig. 1a).

To further characterize the MajSAT RNA expression in pancreatic cancerous cells, we performed northern blotting using a probe containing a single 234 bp array of the MajSAT repeat consensus sequences. Because the transcript lengths of MajSAT RNA were highly heterogeneous, they were detected as smear or ladder bands covering from ~200 to 8,000 nucleotide, which was consistent with other reports\(^18\,26\) (Fig. 1b). In addition, northern blotting using strand-specific probes revealed that MajSAT RNA was transcribed exclusively from the forward strand of the genomic DNA, similarly to human satellite III (satIII) RNA\(^27\), which is expressed under limited circumstances in a strand-specific manner from pericentromeric satellite sequences\(^28\,29\). The expressed MajSAT RNA in cancerous cells was localized mainly to the cytoplasm when testing the fractioned RNAs (Fig. 1c). Interestingly, consistently with the previous report\(^18\,20\), MajSAT RNA expression could not be detected in K512 and K375 cells, established cell lines from PanIN and pancreatic cancer cells in the mice described above, respectively, when they were cultured as a monolayer in a dish. However, MajSAT RNA was re-expressed when the cells were transplanted subcutaneously onto the backs of nude mice as allografts (Fig. 1d).

**MajSAT causes chromosomal and genomic instability.** As described above, MajSAT RNA was expressed in a strand-specific manner in precancerous cells in vivo, but the expression was not observed in tumour-derived cell lines in vitro. Utilizing these results, to characterize the biological effects of the aberrant MajSAT RNA expression in precancerous cells in vitro, we established constitutively MajSAT RNA-expressing cells. We constructed two types of expression constructs: pLV5IN- EF1z-MajSAT, which constitutively expresses forward-strand MajSAT RNA driven by the EF1z promoter, and pTREeight-MajSAT, which expresses forward-strand MajSAT RNA only in the presence of doxycycline (Fig. 2a). Because it is practically difficult to express all lengths of the satellite repeats simultaneously by the currently available methods, approximately three tandem repeats of the basic unit sequences of MajSAT RNA were chosen for expression in this study (Fig. 2a) to examine the effects of the basic sequence and of the junctional sequences between the units. We confirmed the stable expression and the tightly regulated expression by doxycycline, respectively, in the construct transduced cells (Fig. 2b).

Using the NIH3T3 and the PanIN-derived K512 cells stably expressing MajSAT RNA, we first performed a focus formation assay and a soft-agar colony formation assay. In both cases, cells with MajSAT RNA expression showed higher rates of the acquisition of malignant phenotypes, escape from contact-inhibition and acquisition of anchorage-independent growth (Fig. 2c,d), although the cell proliferation rate was not changed by the expression of MajSAT RNA (Supplementary Fig. 1a,b). Similar to previous observations using primary human mammary epithelial cells (HMECs)\(^19\), the frequencies of mitotic errors, such as spindle multipolarity, micronuclei and anaphase bridging, during cell division were significantly increased by
MajSAT RNA expression (Supplementary Fig. 1c–f). While these chromosomal instabilities may lead to losses, gains or translocations of chromosomes, which are observed in pancreatic cancer cells, small mutations at the nucleotide level, such as base substitutions and the insertion or deletion of nucleotides, are crucial in the pathogenesis of PanIN-carcinoma sequence3–5,7. Thus, we next performed comprehensive exome sequencing to examine whether MajSAT RNA expression enhances the spontaneous mutation rate. To equalize the starting mutation levels, K512 cells expressing MajSAT regulated by doxycycline were cultured with or without doxycycline for 4 weeks, exon sequencing was performed to compare the rates of single nucleotide variants (snv) and of small insertions and deletions (indel) in cells with and without MajSAT RNA expression (Supplementary Data 1 and 2; Fig. 2e–g). The results showed that MajSAT RNA expression increased the snv and indel rates (Fig. 2e,f), while the spectra of base alternations were comparable in both cell types, suggesting that more DNA damage (or DNA-damage repair impairment) occurred to the bases universally in cells expressing MajSAT RNA (Fig. 2g).

Moreover, because mtDNA is generally more sensitive and mutagenic under genotoxic stress than is genomic DNA30, we also examined the changes in the mutation rates in mtDNA by MajSAT RNA expression. We sequenced the non-coding D-loop region of mtDNA, which was reported to have a high mutation tendency31. The frequencies of point mutations and small nucleotide insertions/deletions were significantly higher in K512-EF1α-MajSAT cells (Fig. 2h; Supplementary Figs 2 and 3), suggesting that MajSAT RNA significantly disturbs the intracellular homeostasis by enhancing the genomic instability in both nuclear and mtDNA, which may contribute to the pro-carcinogenetic steps in the long-term.

MajSAT RNA is specifically bound to YBX1 protein. To determine the possible molecular mechanisms of the genomic instability induced by MajSAT RNA expression, we searched for proteins interacting with MajSAT RNA, as non-coding RNAs frequently function by interacting with specific proteins32,33. Nuclear and cytoplasmic lysates from K512 cells were mixed with BrU-labelled forward- (as samples) and reverse-strand
Figure 2 | MajSAT RNA causes chromosomal and genomic instability. (a) Two constructs expressing MajSAT RNA used are shown. EF1α promoter-driven constitutively MajSAT RNA-expressing plasmid (upper; EF1α-MajSAT) and tetracycline-regulated MajSAT RNA-expressing plasmid (lower; TREtight-MajSAT), for which the MajSAT RNA expression can be shut off by removing tetracycline. Both constructs express approximately three tandem repeats of the 234 nucleotide basic unit of MajSAT RNA. Dox, doxycycline; Maj, MajSAT basic unit. (b) Confirmation of MajSAT RNA expression in K512 cells stably transfected with EF1α-MajSAT (left) or TREtight-MajSAT (right) by northern blotting. Dox +, doxycycline at 1 μg ml⁻¹ for 48 h. (c) Focus formation assay of NIH3T3 stably MajSAT RNA-expressing (NIH3T3-MajSAT) cells. The number of piled-up foci that escaped from contact growth inhibition was counted after 14 days culture. (d) Soft-agar colony formation assay using K512 stably expressing MajSAT RNA (K512-MajSAT) cells. Representative cell images are shown at left. Bar, 50 μm. Data on the number of foci represent the mean ± s.e. of three independent experiments (right). *P < 0.05 (Welch’s t test). (e, f) MajSAT RNA expression increases genomic instability. Whole-exon sequences in K512-TREtight-MajSAT cells after culturing with or without doxycycline for 4 weeks were examined by high-throughput sequencing. The total numbers of acquired somatic mutations (e) and small insertions/deletions (f) in each group are shown. (g) The spectrum of acquired somatic mutations. The frequencies of the indicated base change patterns are shown. White and black bars indicate cells cultured without and with doxycycline, respectively. (h) Frequencies of mutations in the mtDNA. The cloned libraries of the D-loop region of mtDNA, generated from DNA extracts of K512-vector (vector) and K512-MajSAT (MajSAT) cells, were sequenced. Mutation frequencies out of total sequenced nucleotides are summarized. Data represent the mean ± s.e. of two independent experiments. *P < 0.05 (Student’s t test).

Although YBX1 localizes to the cytoplasm under normal conditions, nuclear translocation was reported under various stress conditions such as oxidative stress, ultraviolet irradiation and the treatment with DNA-damage inducing agents. Several reports showed that translocated YBX1 enhances DNA-damage repair machinery by interacting with DNA-damage repair genes. Because MajSAT RNA is expressed in the cytoplasm, as determined by the northern blot results shown above, and binds to YBX1 in the cytoplasm, we hypothesized that the translocation of YBX1 into the nucleus was inhibited by its trapping in the cytoplasm by MajSAT RNA. First, fluorescence in situ hybridization (FISH) analysis using K512 cells transiently transfected with a MajSAT RNA-expressing...
plasmid (pLVSIN-EF1α-MajSAT) confirmed the primarily cytoplasmic distribution of MajSAT RNA. Probe specificity was confirmed by the disappearance of the targeted RNAs following RNase treatment (Supplementary Fig. 4). Next, to determine the intracellular localization of MajSAT RNA and YBX1 protein, we performed double staining of MajSAT RNA and YBX1 protein by in situ hybridization and immunofluorescence staining using K512 cells transiently transfected with MajSAT RNA-expressing plasmid (pLVSIN-EF1α-MajSAT), which enabled us to compare MajSAT RNA-expressing and non-expressing cells in the same field of view. MajSAT RNA was localized mainly to the cytoplasm, which was consistent with the northern blotting results (Fig. 1c). Although YBX1 was diffusely distributed in the cytoplasm in the cells not transfected with MajSAT RNA, YBX1 was aggregated into particles in the cytoplasm and co-localized with MajSAT RNA in MajSAT RNA-expressing cells even without any other stimulation (Fig. 3d, upper panels). Furthermore, the nuclear translocation of YBX1 under oxidative stress induced by H2O2 treatment was inhibited in MajSAT RNA-expressing cells (Fig. 3d, lower panels). The percentages of nuclear translocation of YBX1 were 6.7 ± 3.6% in MajSAT RNA-expressing cell and 74.5 ± 8.2% in non-expressing cells, respectively (mean ± s.e. from the results of twenty fields observation). These phenomena were similarly observed after DNA-damage induction by ultraviolet irradiation (Supplementary Fig. 5a,b). To confirm the specificity of YBX1 retention by MajSAT RNA, K512 cells expressing reverse-strand MajSAT RNA (MajSAT Rv) were also established as a negative control (Supplementary Fig. 6a). Interestingly, unlike sense MajSAT RNA, antisense MajSAT RNA localized to the nucleus (Supplementary Fig. 6b), consistent with a previous report15, and did not impair YBX1 translocation to the nucleus (Supplementary Fig. 6c). Nuclear translocation of proteins known as YBX1-interacting proteins, RBPp6, CTCF1, ANKRd2 and p53 (ref. 22), was also evaluated following H2O2 treatment. Among the proteins analyzed, RBPp6 translocation to the nucleus after
stress induction was impaired (Supplementary Fig. 7a–d). This result suggests that the impaired nuclear transport of YBX1 by MajSAT RNA could affect the nuclear transport of YBX1-interacting proteins, ultimately affecting their intracellular localization.

**MajSAT RNA impairs DNA-damage repair activity.** To examine whether the increased mutation rates in MajSAT RNA-expressing cells were due to the increased DNA damages by the inhibition of the intra-nucleus YBX1 function, we established MajSAT RNA-expressing K512 cells with overexpression of GFP-tagged YBX1 (K512-MajSAT-YBX1GFP; Supplementary Fig. 8a,b) to overcome the binding of YBX1 to MajSAT RNA in the cytoplasm. The expression levels of YBX1 protein were approximately doubled in these cells (Supplementary Fig. 8a), and the nuclear translocation of YBX1 was rescued by H2O2 treatment was rescued (Fig. 4a). Using these cells, we measured cellular levels of 8-hydroxy-2’-deoxyguanosine (8-OHdG), the most commonly used indicator of oxidative DNA damage, before and after H2O2 treatment. The 8-OHdG levels at 2 and 24 h after H2O2 treatment were significantly increased in MajSAT RNA-expressing cells, whereas the effect was negated at 24 h by YBX1 protein overexpression (Fig. 4b).

Because of this time course and because the cellular reactive oxygen species (ROS) levels were not significantly changed in these cells (Fig. 4c), we hypothesized that the increased 8-OHdG levels in MajSAT RNA were caused by the impairment and retardation of DNA-damage repair. Because 8-OHdG is repaired mainly by the base excision repair (BER) pathway35,36, we determined the BER activity using the colorimetric DNAlzyme-based assay37,38. BER activity in the MajSAT-expressing K512 cells was significantly reduced even in the steady condition. Furthermore, BER activities were enhanced in K512-vector control cells after H2O2 treatment, while such enhancement was hardly observed in MajSAT RNA-expressing cells (Fig. 4d).

To further examine the rescue effects of YBX1 overexpression in MajSAT-expressing cells, we performed the hypoxanthine phosphoribosyl transferase (HPRT) mutation assay. In this assay, once cells acquire HPRT gene missense mutations, they survive and form colonies under selection by 6-thioguanine (6-TG), which is converted into cell-toxic 6-mercaptopurine by functional HPRT. As expected, 6-TG resistant colonies increased in direct

**Figure 4 | MajSAT RNA reduces DNA repair function via YBX1.** (a) MajSAT RNA reduces YBX1 movement into the nucleus after oxidative stress, which is recovered by the forced expression of YBX1. K512 cells stably expressing MajSAT RNA or MajSAT RNA with the forced expression of YBX1 were used. Cells were treated with or without 400 μM H2O2 for 6 h. The nuclear entry of YBX1 was determined by immunofluorescence staining. Representative images from three independent experiments are shown. Bar, 10 μm. (b) The recovery of oxidative DNA damage was delayed in MajSAT RNA-expressing cells (MajSAT), which was countered by the forced expression of YBX1 (MajSAT + YBX1). The levels of 8-OHdG were determined by competitive ELISA after the treatment of cells with 400 μM of H2O2 for the indicated periods. Data represent the mean ± s.e. of three independent experiments. *P < 0.05 (Student’s t test). (c) Cellular ROS levels were not significantly changed by MajSAT RNA or YBX1 expression. Average luminescences, reflecting cellular ROS levels, were measured using K512 cells before and after 400 μM H2O2 stimulation for 2 h. (d) BER activity was lower in MajSAT RNA-expressing cells, which was rescued by YBX1 forced expression. The 8-OHdG excision activity was determined by the colorimetric DNAzyme assay. To determine the response to oxidative stress, cells were incubated in 400 μM H2O2 for 4 h. Data represent the mean ± s.e. of three independent experiments. *P < 0.05, (Student’s t test). (e) Spontaneous missense mutations in the HPRT gene occurred more frequently in MajSAT RNA-expressing K512 cells. Cells were treated with 2 μM MNU for 1 or 2 weeks to promote the tendency for mutation and subsequently cultured in 6-TG-containing media. The number of surviving colonies, in which mutations in the HPRT gene presumably occurred, was counted. Data represent the mean ± s.e. of three independent experiments. *P < 0.05 (Welch’s t test).
proportion to the mutagenic MNU treatment period (Fig. 4e). However, the number of surviving colonies expressing MajSAT RNA was significantly higher and the phenomenon was clearly rescued by YBX1 expression (Fig. 4e).

Finally, we observed the reduced copy numbers of mtDNA (mt-Co1 and mt-Cytb) in MajSAT RNA-expressing cells, reflecting the increased mtDNA mutations\(^{39}\), which were also rescued by YBX1 protein overexpression (Supplementary Fig. 8c). All of these results suggested that MajSAT RNA expression impairs intrinsic YBX1 function and leads to an increase in the number of genomic and mtDNA mutations.

To exclude the possibility that the effects of MajSAT RNA harbouring only three repeats of the basic unit were specific to that particular construct, we generated K512 cells expressing MajSAT RNA with six repeats (Supplementary Fig. 9a) and confirmed elevated 8-OHdG levels, increased numbers of 6-TG resistant colonies and decreased copy numbers of mtDNA as a result of genomic instability (Supplementary Fig. 9b–d). Although this does not reproduce the endogenous heterogeneous conditions, these results suggest that the basic units of MajSAT RNA are likely responsible for the phenotypes caused by MajSAT RNA shown here.

MajSAT RNA-non-interacting mutant recovers YBX1 function.

To further confirm the biological significance of the YBX1-MajSAT RNA interaction, we constructed a series of flag-tagged YBX1 deletion constructs to determine the domain of YBX1 responsible for its interaction with MajSAT RNA (Supplementary Fig. 10a,b). We found that the C terminus (amino acids 218–323) of YBX1 is crucial for its interaction with MajSAT RNA, because the flag-tagged C-terminal YBX1 deletion mutant (hereafter referred to as FmYBX1) no longer interacted with MajSAT RNA, according to RNA immunoprecipitation (Supplementary Fig. 10b). To replace the wildtype YBX1 with FmYBX1 in K512 cells, siRNAs targeting the 5’ untranslated region of YBX1 were transfected into K512 cells stably expressing FmYBX1, resulting in efficient knock down of endogenous wildtype YBX1 but not FmYBX1 (Supplementary Fig. 10c). The nuclear translocation of FmYBX1 following \(H_2O_2\) treatment was retained in these cells (Supplementary Fig. 10d). Furthermore, reduced BER activity, increased 8-OHdG levels, increased 6-TG resistant colonies and decreased mtDNA levels were recovered in FmYBX1- and MajSAT RNA-expressing cells (Supplementary Fig. 10e–h). The results observed with the MajSAT RNA-non-interacting mutant of YBX1 suggest that the MajSAT RNA–YBX1 interaction impairs YBX1 function, resulting in an increased number of genomic and mtDNA mutations.

Discussion

Herein, we show that MajSAT RNA is aberrantly expressed in mouse PanIN (mPanIN) cells carrying a constitutively active K-ras gene, and it increases the number of genomic and mtDNA mutations by binding with YBX1, in addition to increasing the mitotic instability. All of these effects may promote the cellular transformation of precancerous cells.

The regulatory mechanisms of satellite transcription from the centromeric and pericentromeric genomic regions, which are normally in a heterochromatin state, are not yet fully clarified. Although it is suggested that epigenetic alternations in cancer cells such as global DNA hypomethylation may induce aberrant transcription from heterochromatin regions\(^ {26,40,41}\), this hypothesis still remains controversial\(^ {42,43}\). Nonetheless, aberrantly high expression levels of satellite RNAs have been reported in various types of epithelial cancers, including pancreatic cancer and colon cancer, both of which have higher rates of K-ras gene mutations\(^ {18,20}\). Importantly, the aberrant expression of satellite RNAs was also observed in PanIN tissues\(^ {18}\). Consistent with previous reports, we also detected aberrantly high expression levels of MajSAT RNA in mouse PanIN tissues with genetically induced K-ras mutations. Although a few reports have demonstrated possible regulator of satellite transcription, such as Snail1/LOXL2 (ref. 44), p53 (ref. 40) or Pax3 and Pax9 (ref. 45), K-ras mutations are apparently among the potent inducers of aberrant MajSAT RNA transcription while the downstream events are not fully clarified.

Aberrantly highly expressed satellite RNAs in precancerous cells may have oncogenic potential in the sequential carcinogenesis model. Consistent with a previous report in which the loss of BRCA1 in mammalian cancer cells causes derepression of satellite RNAs by H2A monoubiquitination and induces centromere amplification\(^ {19}\), MajSAT RNA overexpression did induce chromosomal instability, such as spindle multipolarity, micronuclei and anaphase bridging. In addition, we determined that MajSAT RNA expression might induce the acquisition of somatic mutations in the genome, possibly resulting in the occurrence of mutations in the defined driver genes during the sequential carcinogenesis process. These results, together with the very recent findings that satellite RNA transcripts stimulate the production of proinflammatory cytokines\(^ {46}\) and that human satellite II (HSATII) transcripts lead to the progressive elongation of pericentromeric regions by intrinsic reverse transcriptases\(^ {20}\), may indicate that aberrant satellite RNA expression plays more crucial roles in long-term carcinogenesis than previously considered.

Another important finding in this study is that the aberrant expression of satellite RNA increases mutations in mtDNA. As recently reported in the *Drosophila* model, epithelial cells that are defective in mitochondrial function in conjunction with oncogenic Ras expression potently induced tumour progression in the surrounding tissues\(^ {47}\). The high rates of mitochondrial malfunction in cancers are caused by somatic mutations in the mtDNA\(^ {48,49}\), which are frequently identified in pancreatic cancers\(^ {50}\) and the accumulation of mtDNA mutations is already apparent in precancerous lesions\(^ {30}\). From these results, the aberrant expression of MajSAT RNA in PanIN cells might drive not only cell-autonomous but also non-autonomous carcinogenesis in the neighbouring cells via these mechanisms.

We identified YBX1 as a binding protein of MajSAT RNA. Although some previous reports demonstrated that satellite transcripts are normally localized to the nucleus\(^ {45,51}\), we found that aberrantly expressed MajSAT RNA is located mainly in the cytoplasm *in vivo* and *in vitro* and impairs the translocation of YBX1 into the nucleus. While intracellular localization of some YBX1-interacting proteins, such as RBBP6, is also affected by MajSAT RNA expression, YBX1 is one of the key factors for genomic DNA-damage repair\(^ {22}\). Various mechanisms for DNA repair by YBX1 have been suggested, such as endonuclease activity\(^ {52,53}\), the transcriptional enhancement of repair genes\(^ {54,55}\), the enhancer of DNA glycosylation to excise damaged nucleotide bases\(^ {24,34}\) and cooperation with p53 in the nucleus\(^ {56}\). In addition, YBX1 is also a key molecule for mtDNA-damage repair, in which fewer numbers of molecules participate compared with genomic DNA repair\(^ {37,58}\). Therefore, it is speculated that the effects of YBX1 dysfunction are more severe in mtDNA repair than genomic DNA repair. This difference may be reflected by our results indicating that the copy number of mitochondrial DNA was significantly lower in MajSAT-expressing cells, most likely due to the rapid removal of deteriorated mitochondria\(^ {39}\), which also plays an active role in the aetiopathology of cancer according to Warburg’s hypothesis\(^ {47}\).
In this study, we found that MajSAT RNA is distributed primarily in the cytoplasm. There are inconsistencies among subcellular distribution analyses; however, several reports have identified satellite RNA in the nucleus [13,20], and others have identified it in both the nucleus and cytoplasm [15,28]. During the developmental stages, nuclear and cytoplasmic distribution of MajSAT RNA has been reported [13,20], and it is noteworthy that only the sense sequences were found in both the cytoplasm and nucleus, while antisense transcripts were found only in the nucleus. Although the reasons for these discrepancies are unknown, it will be important to exclude the possibility of genomic DNA cross-hybridization when MajSAT RNA is observed in the nucleus, since MajSAT RNA is encoded by multiple regions throughout the genome. In addition, differences in methods used for situ hybridization, including fixation, permeabilization and proteinase treatment, may affect the ability to determine the distribution of MajSAT RNA.

We also found that the sense sequences of MajSAT RNA are expressed in pancreatic cancerous tissues. While accumulation of satellite RNAs in both orientations has been observed in other conditions [45,51], previous reports have also shown that strand-specific transcription of major satellite RNA occurs during the 2-cell stage of the mouse embryo [15], and an increase in forward-strand satIII transcription was observed following MajSAT RNA has been reported [15,29], and it is noteworthy that only RNAs, will be necessary to interpret these results fully. Addressing the orientation and intracellular distribution of MajSAT may explain some of the discrepancies. Further experiments addressing the orientation and intracellular distribution of MajSAT RNAs, as well as the expression levels and heterogeneity of satellite RNAs, will be necessary to interpret these results fully.

In this study, we expressed MajSAT RNAs with three or six repeats of the consensus sequence and assessed their biological function. The endogenous MajSAT RNAs, however, are more heterogeneous with additional repeats. While there is currently no adequate method to express all heterogeneous MajSAT RNAs in vitro, our results may suggest that the basic components of the repeat sequences alone have pathological effects even at lower expression levels.

In summary, our findings reveal that aberrantly expressed satellite transcripts may act as ‘intrinsin mutagens’ and have oncogenic potential. Although satellite sequences are not conserved among eukaryotes, and it still remains unclear which sequences in the human genome are the actual counterparts of mouse MajSAT; the sequences HSATII and satIII, which are also located in the pericentromeric region and are aberrantly transcribed in human PanIN tissues [18], may be such candidates [43]. Further elucidating their deregulated expression mechanisms and the downstream events may provide unexpected clues toward the prevention of carcinogenesis.

**Methods**

**Mouse and cell lines.** Tgbr2neo/neo (ref. 60), Ptf1a chooses (ref. 61) and LSL-KrasG12D/+ mouse lines were intercrossed to generate Ptf1aMajSAT. LSL-KrasG12D/−/−; Tgbr2neo/neo (KrasG12D + Tgbr2−/−) and Ptf1aMajSAT. LSL-KrasG12D/−/−; Tgbr2neo/neo (KrasG12D) mice in a > 95% C57BL/6 background. All of the experimental protocols were approved by the internal ethics committee for animal experimentation and conducted in accordance with the Guidelines for the Care and Use of Laboratory Animals of the Graduate School of Medicine, University of Tokyo (Tokyo, Japan).

Mouse primary pancreatic cells derived from mPanIN, which is considered the adenomatous premalignant state of pancreatic tissue, and pancreatic cancer tissues were established from 9-week-old male KrasG12D mouse (K512 cells) and 8-week-old male KrasG12D + Tgbr2−/− mouse (K375 cells), respectively, as previously described [63,64]. Briefly, freshly isolated tumour specimens were cut with sterile razor, digested with dispase II (Invitrogen, Carlsbad, CA, USA)/collagenase (Invitrogen) (4 mg/ml each) for 1 h at 37 °C, and then resuspended in Lowell Park Memorial Institute (RPMI) media supplemented 20% fetal bovine serum and seeded on nitrophen/fibronectin coated plates. After the cells reached pure population, cells were cultured in collagen-coated dishes in RPMI with 10% bovine serum. NIH3T3 and 293TN cell lines were purchased from the American Type Culture Collection (ATCC, Manassas, VA, USA) and System Biosciences (SBI, Mountain View, CA, USA), respectively. These cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM) media supplemented with 10% fetal bovine serum. All cells were incubated at 37 °C, 20% O2 and 5% CO2.

**Subcutaneous allograft model.** For allogenic transplantation, 1 × 105 of K512 and K375 cells in 150 µl of RPMI media was mixed with 100 µl of Matrigel basement membrane matrix, LDEV-Free (Corning, Corning, NY, USA) and injected subcutaneously into the back of 9-week-old female C57BL/6 mice (CLEA Japan, Tokyo, Japan). The resulting tumours were excised after 4 weeks.

**Plasmids.** MajSAT sequences were obtained from a 788 bp array at chromosome 2: 98506702-98507189 (Build: UCSC mm9), which contains almost three tandem repeats of mouse major satellites, as described previously [46]. The polymerase chain reaction (PCR) product was cloned into pCR2.1-TOPO vector (Invitrogen) by the TA cloning method using One Shot DH5α TOP10 competent cells (Invitrogen)

The primers used were as follows: Fw: 5′-CGTTCACAAGATGTGTTT-3′ and Rv: 5′-TGGAAACAGATGTGTTT-3′. The ligated plasmids were named pCR2.1-MajSAT-Fw for the forward-strand insertion and pCR2.1-MajSAT-Rv for the reverse-strand insertion.

The pLV55-EF1a-MajSAT vector was constructed as follows. MajSAT sequences in the pCR2.1-MajSAT-Rv/Fw vector were digested with EcoRI restriction enzyme, subcloned into the pCDH-CMV-MCS-EF1-Puro vector (SBI) (named as pCDH-MajSAT-Fw/Rv vector) and subsequently subcloned into the pLV55-CDH-Puro vector (TaKaRa, Shiga, Japan) by digestion as the XbaI and NotI sites. To append the polyA signal downstream of the MajSAT sequences, SV40 polyA signal sequences from pCDH-CMV-MCS-EF1-Puro vector at the positions of 4534–5656 were amplified by PCR and cloned into the pLV55-CDH-Puro vector at the NotI site using the InFusion HD cloning kit (Clontech, Mountain View, CA, USA) according to the manufacturer’s protocol (named as pLV55-MajSAT-Fw/Rv vector). The primers used were as follows: Fw: 5′-ATCCGTTAAATTCGCTAGCTCTAGCGAATTCGCCCTTTGGAAA-3′ and Rv: 5′-AGATCCATGGGCGCGCCAATATTGATGTGTTT-3′.

The lentiviral particles were named pCR2.1-MajSAT-Fw vector into the pTREtight vector (Clontech) at the EcoRI site. The pB-LCMV-MajSAT-Fw/Rv vector was constructed using the pB-LCMV-EF1 vector (Clontech). MajSAT Fw sequences digested with EcoRI and EcoRV were digested into the multiple cloning site (MCS) and MajSAT Rv sequences digested with BgIII were subcloned into the MCSII site of the pBLCMV vector.

To double the number of MajSAT unit sequences (MajSAT-double), the pCR2.1-MajSAT-Fw/Rv vector was used to transfer into 293TN cells using Effectene Transfection Reagent (Qiagen, Hilden, WI, USA). Briefly, 24 h after 1 g of pPACKH1 packaging plasmid mix were FuGENE HD Transfection Reagent (Promega, Madison, WI, USA). Briefly, 24 h after 1 × 105 cells were seeded in a 6-well dish, 1 µg of template vector and 3 µl FuGENE Reagent diluted in serum-free media were added. To generate stably expressing polyclonal cells, Lentivirus Packaging System (SBI) was used according to the manufacturer’s protocol. Briefly, 1 µg of MajSAT- or YBX1-overexpressing vector and 5 µg of pPACKH1 packaging plasmid mix were transfected into 293TN cells using Effectene Transfection Reagent (Qiagen, Hilden, Germany). After 24 h, the collected culture media was mixed with one-fifth the volume of PEG–it Reagent (SBI) overnight at 4 °C to concentrate the virus. The centrifuged pellet was resuspended in 1 × PBS, and aliquots were stored at −80 °C. The tissues were homogenized with 20 µl of lysis buffer and sonicated for 7 × 30 s each on ice. The lysates were added to the targets with Polybrene Reagent (Santa Cruz Biotechnology, Dallas, TX, USA) and after 48 h, antibiotic selection (3 µM puromycin or 150 µM neomycin) was begun.

**Transfection and lentivirus transduction.** Transient transfection of the plasmids was performed using FuGENE HD Transfection Reagent (Promega, Madison, WI, USA). Briefly, 24 h after 1 × 105 cells were seeded in a 6-well dish, 1 µg of template vector and 3 µl FuGENE Reagent diluted in serum-free media were added. To generate stably expressing polyclonal cells, Lentivirus Packaging System (SBI) was used according to the manufacturer’s protocol. Briefly, 1 µg of MajSAT- or YBX1-overexpressing vector and 5 µg of pPACKH1 packaging plasmid mix were transactivated into 293TN cells using Effectene Transfection Reagent (Qiagen, Hilden, Germany). After 24 h, the collected culture media was mixed with one-fifth the volume of PEG–it Reagent (SBI) overnight at 4 °C to concentrate the virus. The centrifuged pellet was resuspended in 1 × PBS, and aliquots were stored at −80 °C. The tissues were homogenized with 20 µl of lysis buffer and sonicated for 7 × 30 s each on ice. The lysates were added to the targets with Polybrene Reagent (Santa Cruz Biotechnology, Dallas, TX, USA) and after 48 h, antibiotic selection (3 µM puromycin or 150 µM neomycin) was begun.
To generate Tet-induced MajSAT-expressing cells, K512 cells were first transfected with the pTet-On-advance vector (Clontech) following 150 μM neomycin selection. Tet-ON-advanced cell clones were subsequently transfected with the pTRE-edit-MajSAT vector along with the Linear Hygromycin Marker (Clontech). MajSAT RNA expression after induction by doxycycline (1 μg·mL⁻¹) was confirmed in the clones selected with 150 μM hygromycin.

For transcription of sIRNAS targeting X1R2, the pre-designed TriFecta Dicer-Substrate RNAi Kit for mouse X1R2 (Integrated DNA Technologies, Coralville, IA, USA) was purchased, and 5 × 10⁴ cells were transfected with 10 nmol of each siRNA oligo in 12-well plates using Lipofectamine RNAiMAX Transfection reagent (Thermo Fisher, Waltham, MA, USA) according to the manufacturer's protocol. The sIRNA sequence of RNAiF: 5'-UUGUCCAUACCCAAAGAAGAAUGAGGAGCAGGACCAAGGGTC-3'.

Probe synthesis. For northern blotting and RNA in situ hybridization of mouse pancreatic tissues, transcription of pCR2.1-MajSAT-cDNA vectors by transfection using MEGAscript T7 Kit (Ambion, Austin, TX, USA), followed by biotin labelling using the Psoralen-Biotin Linker Kit (Roche Diagnostics, Basel, Switzerland). Briefly, slides were blocked by a quick sonication. Homogenized samples were added to 700 μL of NE buffer followed by centrifugation at 16,000 g for 20 min. After supernatant were strongly denatured at 55 °C in 2 × SSC containing 0.1% SDS and in 0.1 × SSC containing 0.1% SDS, twice each, and the bound probe was visualized using a BrightStar BioDetect Kit (Ambion), according to the manufacturer's protocol.

Focus formation and soft-agar colony formation assay. To examine their ability to escape from contact-inhibition, NIH3T3 cells on a 10 cm dish were maintained in confluency for 2 weeks, changing the media every two days. Piled-up foci were counted by crystal violet staining. To evaluate the acquisition of anchorage-independent growth, 1.5 × 10⁵ of K512 cells were added to 1.5 ml of plating agar, consisting of 1× RPMI and 10% FBS containing 0.35% agarose (Sigma-Aldrich), and poured onto a base RPMI agar containing 0.5% agarose in a 6-well dish. Colonies grown to over 50 μm in size were counted after 3 weeks of incubation.

Exome sequencing. K512-TRE-edit-MajSAT cells were cultured in RPMI media with and without 1 μg·mL⁻¹ of doxycycline and passaged every 3 days for 4 weeks. DNA was extracted using the QIAamp DNA mini kit (Qiagen). Three micrograms of qualified genomic DNA was fragmented by an ultrasonicator Covaris S-series (Covaris, Woburn, MA, USA). The fragments were purified, end-blunted, A-tailed, and adaptor ligated using the Agilent SureSelectXT Human All Exon v4 Kit (Agilent Technologies, Santa Clara, CA, USA) in accordance with the manufacturer's protocol. sIRNA sequencing Target Enrichment for Illumina Multiplexed Sequencing Protocol. Five cycles of PCR were performed after size selection in the gel. Then, 750 ng aliquots of these purified libraries were hybridized to the SureSelect oligo probe capture library (Agilent) for 24 h. After hybridization, washing and elution with magnetic beads, the eluted fraction was PCR-amplified for 12 cycles with primer pairs, quantified using qPCR and an Agilent 2100 Bioanalyzer (Agilent) to obtain sufficient DNA template for the downstream applications. Paired end sequencing was performed by the HiSeq 2000 according to the manufacturer’s instructions (Illumina, San Diego, CA, USA). After preprocessing reads containing sequencing adapters and low-quality reads with more than five ambiguous bases, high-quality reads were aligned to the UCSC mouse reference genome (mm9) using BWA (v0.5.9) with default parameters. Picard (v1.7.0) (http://picard.sourceforge.net/) was used to mark duplicates. Somatic point mutations and somatic indels were detected by VarScan2.3.2 and SAMtools (http://samtools.sourceforge.net/) with default parameters. ‘Somatic’ mutation was defined as mutations acquired in MajSAT RNA-expressing cells. ‘LOH’ was defined as mutations acquired in cells not expressing MajSAT RNA. Detected snv and indel mutations were annotated using the snpEff tool (v3.0) [68].

Northern blotting. Northern blotting was performed as described previously with slight modification. Briefly, 5 μg of tissue RNA was separated in 1% formaldehyde denatured agarose gel and hydrostically transferred to Hybond N+ membrane (GE Healthcare, Chalfont St Giles, UK). Membranes were ultraviolet-crosslinked and hybridized in the hybridization buffer. Hybridization was performed overnight at 42 °C in ULTRAhyb Buffer (Ambion) containing 10 ng·mL⁻¹ of biotin-labelled RNA probe, which had been denatured at 90 °C for 10 min. After prehybridization, the blots were strained with washing buffer (T2, 0.1× SSC, 50% formamide) were applied for washing twice with 1× PBS, antigens were degraded with 70, 95 and 100% ethanol. Next, 50 ng·mL⁻¹ of biotin-labelled MST-Rv probe, 500 μg·mL⁻¹ of yeast tRNA and 100 μg·mL⁻¹ salmon DNA in the hybridization buffer (5× SSC, 50% formamide) were applied to the slide glass and incubated overnight at 50 °C. After stringent washing with 5× SSC for 5 min, 1× SSC for 5 min and 0.1× SSC for 5 min at 50 °C, the slides were blocked and stained using the DIG Wash & Block Kit (Roche Diagnostics, Basel, Switzerland). Briefly, slides were blocked for 30 min with 1× DIG block buffer and incubated with 0.2 μg·mL⁻¹ Streptavidin-Alkaline Phosphatase (Ambion) diluted in 1× block buffer for 30 min. After washing three times with 1× wash buffer, the targets were then visualized using NBT/BCIP solution (Sigma-Aldrich, St Louis, MO, USA) with levamisole (Dako, Glostrup, Denmark) for 10 min and counterstained with Nuclear Fast Red (Sigma-Aldrich) for 1 min.

RNA extraction. Mouse pancreatic tissue was immediately frozen in liquid nitrogen after resection and stored at −80 °C. Frozen tissues were crushed without thawing by a homogenizer (Tokyo Rikakiki, Japan) and immediately immersed in ice-cold Isogen reagent (Nippon gene, Tokyo, Japan). Nuclear and cytoplasmic RNA were isolated separately using the Cytoplasmic & Nuclear RNA Lysis Kit (Ambion) for 30 min and subsequently mixed with 1× Proteinase K and incubated at 65 °C for 30 min. The total RNA sample was then precipitated using isopropanol for 30 min at −20 °C. Washing with 70% ethanol and drying in air followed by measurement of the absorbance at 260 nm using a microplate reader (Bio-Rad Laboratories, Hercules, CA, USA).

Cell growth. For the cell growth assay, the Cell Counting Kit-8 (Dojindo Molecular Technologies, Kumamoto, Japan) was used according to the manufacturer’s protocol. Briefly, 1 × 10⁶ cells were seeded in 6-well plates. On the next day, 104 cells were seeded in 96-well plates. Then, 10 μL of Cell Counting Kit-8 solution was added, and the plates were incubated for 90 min at 37 °C, followed by measurement of the absorbance at 450 nm using a microplate reader (Bio-Rad Laboratories, Hercules, CA, USA).

mtDNA mutation. To examine the mutation rate in the mitochondrial DNA, a 431 bp fragment of the D-loop region was cloned using the InFusion HD cloning kit (Clontech) as follows. Total DNA templates including mtDNA were isolated from K512-EF1α-MajSAT cells using the QIAamp DNA Mini Kit (Qiagen), and PCR was performed using LA-Taq polymerase (TaKaRa) mixed with Pfu turbo DNA Polymerase (Stratagene). Primers used were as follows: Fw: 5’-CTGCACTTGGATCCTCTTTTTATTTTGGCCTGATCTTT-3’ and Rev: 5’-AACGAGTCGACGATCCATCCTATAGGCTTCCCTGG-3’. The PCR product was digested at the BamHI site and dephosphorylated with Shrimp Alkaline Phosphatase (TaKaRa) at 37 °C for 15 min. Amplified fragments and linearized vector were incubated with 5 × InFusion HD Enzyme premix at 50 °C for 15 min and subsequently transformed into ECOS Competent E. coli DH5α. After 16 h, at least twenty colonies from each sample were picked up, and restricted vectors were collected using the QIAprep 96 Turbo Miniprep Kit (Qiagen). The inserted D-loop sequences of the cloned libraries were determined using a 72 nucleotide primer (5’-TAA TAC GAC TCA TAT GGT G-3’). The mutations were determined by comparison with the reference sequences (GenBank: NC_005089.1).
ion trap mass spectrometer (LC-MS/MS) with a slight modification. The individual spectra from MS/MS were processed using the TurboSEQUEST software (ThermoFinnigan, CA, USA). The generated peak list files were used to query either the MSDB database or NCBI using the MASCOT programme (http://www.matrixscience.com).

To examine the endogenous binding of YBX1 protein to MajSAT RNA, the RIP assay microRNA (mRNA) (MBI) was used according to the manufacturer’s protocol. Briefly, 6 × 10⁵ K512-HA-YBX1 cells were seeded, and 8 μg of BrU-labelled MajSAT-Fw RNA was transfected into the cells using TransMessenger reagent (Qiagen). After 8 h incubation, cells were harvested and lysed, followed by the addition of precleared Protein G magnetic beads for 1 h. The supernatant containing argo beads conjugated to anti-HA tag-antibody or control rabbit IgG for 4 h at 4°C. The pellets were washed four times, and bound RNA was isolated by ethanol precipitation. To detect bound MajSAT RNA, 1 μg of precipitated RNA and 5 μl of input were reverse transcribed to cDNA using Superscript III Reverse Transciptase (Invitrogen), and semi-quantitative RT-PCR was performed using Mighty Amp DNA polymerase (TaKaRa).

**Western blot analysis and antibodies.** Western blotting was performed as previously described69. Briefly, lysate samples were separated on 5-20% gradient polyacrylamide gel by SDS-PAGE following electric transfer to PVDF membranes (GE Healthcare). After blotting with 5% dry milk, membranes were probed with the respective primary antibodies that were detected in immunoblotting (1:1000) in the primary antibody solution (Cosmo Bio, Tokyo, Japan) overnight at 4°C. HRP-conjugated corresponding secondary antibodies (GE Healthcare) were subsequently used. Trueblot anti-rabbit IgG HRP (Rockland, Limerick, PA, USA; 1:1,000) was used to detect the antibody without interfering immunoprecipitated immunoglobulin heavy and light chains.

**Immunofluorescence imaging.** Cells were seeded in 6-well glass chambers (IWAKI) with manual collagen coating. Cells were fixed with 4% PFA in 1× PBS for 15 min at RT and permeabilized with 0.1% Triton-X in 1× PBS for 20 min at room temperature. Primary antibodies diluted by 1:100 with Can Get Signal Immunostaining Solution A (Toyobo, Osaka, Japan) were applied to the samples and incubated at room temperature overnight. Cells were incubated with the secondary antibodies conjugated with Alexa 555 or 488 (Molecular Probe, Eugene, OR, USA) for 1 h for room temperature and mounted using fluorescent mounting medium. The average threshold cycle number (Ct) values of mtDNA (mt-Co1 and mt-Cytb) and nuclear DNA (H19) were obtained, and the relative content of mtDNA, normalized by the Ct values of nuclear-H19, was calculated using the delta-delta Ct method (StepOnePlus Realtime PCR system; Invitrogen). Each sample was run in triplicate.

**RNA in situ hybridization of cultured cells.** The intracellular localization of MajSAT RNA was visualized using the RNA Scope Fluorescent Multiplex Reagent (Agilent) and nuclear DNA (H19) were obtained, and the relative content of mtDNA, normalized by the Ct values of nuclear-H19, was calculated using the delta-delta Ct method (StepOnePlus Realtime PCR system; Invitrogen). Each sample was run in triplicate.

**mRNA copy number measurement.** Total RNA was isolated from 1 × 10⁶ cells using the QiAamp DNA mini kit (Qiagen) according to the manufacturer’s protocol. Then, 100 ng of template was subjected to quantitative SYBR green PCR (StepOnePlus Realtime PCR system; Invitrogen).

**Cellular ROS measurement.** Cellular ROS levels were measured using the ROS-Glo H₂O₂ assay kit (Promega). Briefly, 1 × 10⁵ cells were cultured in 80 μl of medium on 96-well plates. Then, 20 μl of the H₂O₂ substrate solution was added to the wells, and incubated without 400 μl of H₂O₂ solution. Subsequently, 100 μl of ROS-Glo Detection Solution was added and incubated for 20 min at RT, followed by the measurement of relative luminescence units using a GloMax 96 Microplate Luminometer (Promega).

**In vitro 8-OHdG excision activity assay.** The base excision activity colorimetric assay was performed as previously described with slight modification37,39. This assay is based on an HRP-mimicking DNAzyme consisting of a guanine quadruplex sequence and lambda exonuclease. Endogenous DNA glycosylase excise the 8-OHdG site and produce a 5’ phosphate and a 3’-phospho-unsaturated aldehyde in annealed oligonucleotides. The lambda exonuclease digests the 5’-phosphoryl strand of the annealed DNA to release the complementary DNAzyme sequence, which acts as an HRP, catalyzing the H₂O₂-mediated oxidation of 2,2’-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS²⁺ -Abcam) to generate a colorimetric signal. In this study, 10 μM of 8-OHdG including oligonucleotide (DNA-1), its complementary sequence containing the DNAzyme part (DNA-2) and 5 μl NEBuffer2 (New England BioLabs, Tokyo, Japan) was denatured for 5 min at 90°C, followed by annealing with slow cooling to RT. As a negative control, DNA 3 was used in place of DNA 1. For this experiment, 6 × 10⁵ cells on 10 cm dishes were harvested and incubated in 150 μl of extraction buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl, 0.05% NP-40, 2 mM EDTA, 0.1% PMFS) for 30 min at 4°C, followed by centrifugation at 20,000g for 30 min. Then, 10 μl of the supernatant lyes, 10 μg of lambda exonuclease (Promega) and 20 μl of 8-OHdG was added to the annealed oligonucleotide and incubated for 30 min at 37°C to cleave the 8-OHdG site and digest the DNA 1 strand. Subsequently, 15 μl of 5 μM hemin, 75 μl of 2× HEPES solution (25 mM HEPES, 200 μM NaCl, 10 mM KCl, 0.05% triton, pH 5.2) and 10 μl water were added and incubated for 30 min at RT. Absorption spectra were recorded at 410 nm every 1 min after the addition of 20 μl of 20 mM ABTS²⁺ and 20 μl of 10 mM H₂O₂. The oligonucleotides sequences were as follows: DNA 1: 5’-TCTGCATGGCAATCTTCAAGCGGGCCCTACCC3’ (modified 8-OHdG); DNA 2: 5’-GGGTAGGGCGGTTGGGGGCTGAGAA DNA 3: 5’-TCTGGATCCCCAAGCAGCCTTACCC3’.-1

**MNU-HPR colony formation assay.** K512-vector, K512-MajSAT and K512-MajSAT-YBX1-GFP cells were treated for 1 week with hypoxanthine–aminopterin–thymidine (HAT) medium (Sigma) to eliminate pre-existing mutated cells in the HPRT gene. To promote the opportunities for acquiring oxidation-induced mutations, MNU (Wako) was added for 1 week or 2 weeks. For this experiment, 6 × 10⁵ cells were plated on each 10 cm dish with 5 μM 6-TG (Wako)-containing media, which is catalyzed to cell-toxic 6-mercaptopurine by functioning HPRT. After 20 days, surviving colonies, with potential mutations in the HPRT gene, were counted by violet crystal staining.

**Determination of the YBX1-binding domain.** To construct serial YBX1 deletion mutants (A, B and C), YBX1 coding sequences with various deletions were subcloned into the pLSVIN-EF1 neo vector at the XbaI and BamHI sites using the Infusion HD cloning kit, with a flag tag added to the 5’ end. The pLSVIN-EF1-YBX1GFP vector was used as a template, and the primers used were as follows: Fw: 5’-CCAAATCTCTTACACGACCT-3’ and Rv: 5’-GGCTCATATGCTTGCTGAGG-3’; Ctb-Y; Fw: 5’-TCTGGATTGCAGCATTCATTTCT-3’ and Rv: 5’-AGAGGGAGGTATTTCTTCTGCTC-3’; and mutant YBX1-C; Rv: 5’-GTCACAGGACCAATGATC-3’.

To examine the interaction between the mutant YBX1 protein and MajSAT RNA, DNA immunoprecipitation was performed. Briefly, 1 × 10⁶ 293TN cells were seeded in a 10 cm dish, transfected with the flag-tagged YBX1-overexpression mutant vector or the control vector, and incubated for 48 h.
Cells were harvested, lysed and incubated with precleared Protein G agarose beads for 1 h. The supernatants were mixed with 2 μg MaSat-Fw RNA, which was synthesized by in vitro transcription, and agarose beads conjugated to anti-fla antibody or control IgG for 3 h at 4 °C. The pellets were washed four times, and bound RNA was isolated by ethanol precipitation and resuspended in 10 μl RNase-free water. Precipitation of flange-tagged protein was confirmed by western blotting. To detect bound MaSat RNA, 6 μl precipitated RNA and 5% input were reverse transcribed into cDNA using SuperScript III Reverse Transcriptase (Invitrogen), and semi-quantitative RT-PCR was performed as described earlier. Precipitated samples that were not reverse transcribed were used as a negative control.

Statistical analysis. Statistically significant differences between groups were identified using Student’s t test when the variances were equal. When the variances were unequal, Welch’s t test was used instead. P values <0.05 were considered to indicate statistical significance.

Data availability. The exome sequence data have been deposited in the Sequence Read Archive database (SRA, http://www.ncbi.nlm.nih.gov/sra) under the accession code #SRP081008. All the other data supporting the findings of this study are available within the article and its Supplementary Information and from the corresponding author upon reasonable request.

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