Multiple cancer types demonstrate abnormal expression of repetitive RNA sequences as a form of epigenetic instability. There is growing interest in understanding the role of repetitive RNAs in cancer pathogenesis and immunogenicity and in their potential role as diagnostic or therapeutic biomarkers. In this issue of the JCI, Porter and colleagues report on satellite RNA in a subset of ovarian cancers. The authors found that high expression of human satellite (HSAT) repeats — but not other families of repeats — was associated with an immunosuppressive phenotype in ovarian cancer cell lines and tumor samples. Further induction of HSAT RNA levels in vitro, surprisingly, leads to innate immune activation, suggesting a potential therapeutic strategy. This work highlights the expanding role of repetitive RNAs in tumor biology and the need to better define specific classes of repetitive elements expressed in cancer — as well as their role in tumorigenesis, tumor immunity, and the host response to cancer.
Multiple cancer types demonstrate abnormal expression of repetitive RNA sequences as a form of epigenetic instability. There is growing interest in understanding the role of repetitive RNAs in cancer pathogenesis and immunogenicity and in their potential role as diagnostic or therapeutic biomarkers. In this issue of the JCI, Porter and colleagues report on satellite RNA in a subset of ovarian cancers. The authors found that high expression of human satellite (HSAT) repeats — but not other families of repeats — was associated with an immunosuppressive phenotype in ovarian cancer cell lines and tumor samples. Further induction of HSAT RNA levels in vitro, surprisingly, leads to innate immune activation, suggesting a potential therapeutic strategy. This work highlights the expanding role of repetitive RNAs in tumor biology and the need to better define specific classes of repetitive elements expressed in cancer — as well as their role in tumorigenesis, tumor immunity, and the host response to cancer.

**Repetitive elements in the genome**

Repetitive elements constitute a substantial proportion (more than 50%) of the human genome and include interspersed repeats and tandem repeats (1, 2). Interspersed repeats include the transposable elements (TE), such as short and long interspersed retrotransposable elements (SINE and LINE), and the long terminal repeat-containing human endogenous retroviruses (HERVs). Most transposable elements are stably silenced by DNA methylation and histone silencing marks. The tandem repeats include satellite repeats, microsatellite repeats and telomeric repeats, and are often involved in structural organization of chromosomes and found at or near centromeres and telomeres. α-Satellite repeats are organized in large arrays at all human centromeres. They can produce transcripts in normal cells that are localized in cis to centromeric regions and may be required for efficient loading of centromeric proteins and assembly of functional centromeres. Human satellite II (HSATII) is a satellite repeat family that is found in peri-centromeric regions of a subset of human chromosomes, including chromosomes 1 and 16, which have especially large repeat regions. These HSATII satellitess are usually transcriptionally repressed and organized into constitutive heterochromatin.

**Repetitive RNAs in cancers**

Abnormal expression of repetitive RNA, include TE and satellite repeats, have been reported in multiple cancers, often associated with global DNA demethylation, and may represent a form of epigenetic instability associated with tumorigenesis (5, 6). Repetitive RNA can form double-stranded RNA structures through internal repeats, which can be sensed in the cytoplasm by innate immune pathways evolved to detect viral RNA. These pathways include Toll-like receptors, RIG1, and other proteins, which, upon sensing dsRNA can activate interferon signaling (7, 8). Some transposable elements have intact open reading frames that can translate into an immunogenic protein. Abnormal expression of repetitive RNAs in cancers have been associated with evidence of innate immune activation and interferon signaling (9, 10). Pharmacologic approaches that increase the expression of repetitive elements, including DNA methylation inhibitors and histone modification inhibitors, have been proposed to activate local immune reactions in cancer, potentially increasing intrinsic cancer immunogenicity and enhancing responses to immune checkpoint inhibitors (11,12). There is growing interest in understanding the biological consequences of acquired epigenetic instability due to repetitive elements in cancer, both for diagnostic and therapeutic approaches.

**Satellite RNA in epithelial ovarian cancers**

Porter and colleagues report in this issue of the JCI that a subset of epithelial ovarian cancers (EOCs) express high levels of satellite RNA, which was associated with an immunosuppressive phenotype (13). The investigators performed total RNA sequencing in a set of human cancer cell lines and identified a subset of ovarian cancer cell lines with high expression of satellite elements, including HSATII. High satellite RNA expression, even when present in cell lines that also expressed high levels of other repetitive elements families.
including HERVs, was positively correlated with gene signature of epithelial mesenchymal transition (EMT) and anticoorrelated with interferon (IFN) response pathways. In contrast, cell lines that expressed HERVs and other TEs, but lacked HSATII expression, showed evidence of robust interferon pathway activation.

These findings suggest that HSATII and other satellite repeat expression may play a key role in modulating immune activation in cancer. The association of HSATII expression with EMT and its anti-correlation with IFN was confirmed in a series of EOC samples. Further, analysis of HSATII expression by RNA in situ hybridization in a small cohort of patients with EOCs demonstrated that high HSATII expression was associated with poor outcome. The investigators also showed that extracellular vesicles (EVs) derived from EOC cell lines contained high levels of HSATII RNA as well as other classes of repeat RNA, and suggested that these EVs may deliver HSATII RNAs to other cells, including immune cells, in the tumor microenvironment. Treatment of cell lines with DNA methylation inhibitors induced interferon signaling and expression of TE, including ERVs, LINEs, and SINEs, but not SAT elements. Conversely, treatment with histone-deacetylase inhibitors in multiple cell lines could induce SAT expression but did not strongly induce interferon signalling (13). These findings suggest that manipulating DNA methylation and histone acetylation may have differential effects on different families of repetitive elements. Transfection of locked-nucleic acids targeting HSATII into cell lines with high baseline HSATII expression, paradoxically, further increased HSATII RNA expression and triggered an IFN response, increased MHC1 protein expression, and decreased cell growth (13). This finding suggests that further induction of HSATII expression in HSATII-expressing tumors may break the immunosuppressive phenotype and induce immunogenicity. As Porter et al. have previously identified HSATII expression in pancreatic and other cancers, this finding may have relevance to multiple cancer types (14, 15).

Conclusions
The observations by Porter and colleagues are provocative and demonstrate a complex relationship between repetitive HSATII expression and immunophenotype in ovarian cancer that will require further work to unravel (13). In particular, several important questions are raised. First, what is the mechanism of elevated SAT and HSATII RNA expression in ovarian cancer? The authors posit a role for p53 mutation in the induction of SAT RNA expression (16). However almost all high-grade ovarian cancers have mutant TP53, but only a subset have HSATII expression, suggesting that there are other factors involved, or that only specific mutant alleles of TP53 are functional. The EOC genome is characterized by many structural variations and rearrangements, and the relationship between genomic instability seen in these cancers and the epigenetic instability related to repeat silencing is unclear. Does abnormal expression of SAT RNA disrupt centromere function and contribute to genomic instability, or does underlying genomic instability of these cancers disrupt normal chromatin architecture and contribute to abnormal expression of satellite RNA? In the case of BRCA1, loss of the gene induces structural genomic chaos due to defects in DNA repair but also induces derepression of satellite repeats and disruption of centromere function (17, 18). It would be informative to see if expression of HSATII in EOC and other cancers is associated with specific patterns of chromosomal rearrangements or specific defects in BRCA1-related DNA repair pathways.

It is also puzzling why the expression of satellite RNA, as opposed to that of TE RNA, leads to an immunosuppressive phenotype associated with EMT instead of triggering an IFN response. The nucleus of some cancer cell lines can retain abnormally expressed HSATII transcripts in large foci where they form complexes with MeCP2 and other chromatin regulatory factors, leading to reorganized distribution of polycomb-related protein in the nucleus (5). This sequestering of RNA in the nucleus may prevent RNA sensing systems from recognizing repeat RNAs in the cytoplasm. Similarly, the redistribution of key polycomb-associated genes could affect transcriptional programs and contribute to the EMT phenotype associated with HSATII expression.

The finding by Porter and colleagues highlights the growing importance of acquired epigenetic instability in cancers and demonstrates the complex biology associated with different classes of repetitive elements and the consequences of their dysregulation in cancer (13). Methods to more robustly identify, categorize, and measure expression of different classes of repetitive elements will be needed to further define their roles in tumorigenesis, tumor immunity, and the host response to cancer.

Acknowledgments
SG has received funding from the National Cancer Institute (NCI) and the Department of Defense (DoD).

Address correspondence to: Shridar Ganesan, 195 Little Albany Street, New Brunswick, New Jersey 08901, USA. Phone: 732.235.5211; Email: ganesash@cinj.rutgers.edu.

1. Lander ES, et al. Initial sequencing and analysis of the human genome. Nature. 2001;409(6822):860–921.
2. Richard GF, et al. Comparative genomics and molecular dynamics of DNA repeats in eukaryotes. Microbiol Mol Biol Rev. 2008;72(4):686-727.
3. Chan FL, et al. Active transcription and essential role of RNA polymerase II at the centromere during mitosis. Proc Natl Acad Sci U S A. 2012;109(6):1979–1984.
4. Perea-Resa C, Blower MD. Centromere biology: transcription goes on stage. Mol Cell Biol. 2018;38(18):e00263-18.
5. Hall LL, et al. Demethylated HSATII DNA and HSATII RNA Foci sequester PRCI and Mc2D2 into cancer-specific nuclear bodies. Cell Rep. 2017;18(12):2943–2956.
6. Pappalardo XG, Barra V. Losing DNA methylaton at repetitive elements and breaking bad. Epigenetics Chromatin. 2021;14(1):25.
7. Roers A, et al. Recognition of endogenous nucleic acids by the innate immune system. Immunity. 2016;44(4):739–754.
8. Ori D, et al. Cytosolic nucleic acid sensors and innate immune regulation. Int Rev Immunol. 2017;36(2):74–88.
9. Rooney MS, et al. Molecular and genetic properties of tumors associated with local immune cytolytic activity. Cell. 2015;160(1–2):48–61.
10. Chen R, et al. Endogenous retroelements and the viral mimicry response in cancer therapy and cellular homeostasis. Cancer Discov. 2021;11(11):2707-2725.
11. Vitello GAF, et al. Antiviral responses in cancer: boosting antitumor immunity through activation of interferon pathway in the tumor microenvironment. Front Immunol. 2021;12:782852.
12. Barrero MJ. Epigenetic regulation of the non-coding genome: opportunities for immunoncology. Epigeneomes. 2020;4(3):22.
13. Porter R, et al. Satellite repeat RNA expression in...
epithelial ovarian cancer associates with a tumor immunosuppressive phenotype. *J Clin Invest.* 2022;132(16):e155931.

14. Ting DT, et al. Aberrant overexpression of satellite repeats in pancreatic and other epithelial cancers. *Science.* 2011;331(6017):593–596.

15. Solovyov A, et al. Global cancer transcriptome quantifies repeat element polarization between immunotherapy responsive and T cell suppressive classes. *Cell Rep.* 2018;23(2):512–521.

16. Levine AJ, et al. P53 and the defenses against genome instability caused by transposons and repetitive elements. *Bioessays.* 2016;38(6):508–513.

17. Zhu Q, et al. BRCA1 tumour suppression occurs via heterochromatin-mediated silencing. *Nature.* 2011;477(7363):179–184.

18. Pageau GJ, Lawrence JB. BRCA1 foci in normal S-phase nuclei are linked to interphase centromeres and replication of pericentric heterochromatin. *J Cell Biol.* 2006;173(5):693–701.