Antigenic drift as a mechanism for tumor evasion of destruction by cytolytic T lymphocytes

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It is established that mutations in viral antigenic epitopes, or antigenic drifts, allow viruses to escape recognition by both Ab’s and T lymphocytes. It is unclear, however, whether tumor cells can escape immune recognition via antigenic drift. Here we show that adoptive therapy with both monoclonal and polyclonal transgenic CTLs, specific for a natural tumor antigen, P1A, selects for multiple mutations in the P1A antigenic epitope. These mutations severely diminish T cell recognition of the tumor antigen by a variety of mechanisms, including modulation of MHC:peptide interaction and TCR binding to MHC:peptide complex. These results provide the first evidence for tumor evasion of T cell recognition by antigenic drift, and thus have important implications for the strategy of tumor immunotherapy.


Introduction
It is well established that both neutralizing Ab’s (1, 2) and cytotoxic T lymphocytes can select for viral escape variants in vivo (3, 4). This is usually achieved by mutations that result in replacement of one or more AAs within the antigenic epitopes, a process known as antigenic drift. A recent study revealed that virus-specific CD4 T cells can also select for viral escape variants in vivo (5). Much like viruses, tumors can also evade cytolytic T cells in vivo (6). The described mechanisms include immune ignorance (7, 8), induction of clonal anergy of tumor-specific T cells (9), downregulation of antigen presentation (10, 11), and loss of expression of tumor antigen (12). To our knowledge, it is still unclear whether mutations in tumor antigenic epitopes contribute to tumor evasion of the immune system in vivo.

Transgenic mice expressing T cell receptors for a single antigenic epitope play an instrumental role in establishing antigenic mutation as a mechanism for viral escape of T cell recognition (3, 5). We have recently produced a transgenic mouse line expressing a TCR specific for tumor antigen P1A35-43 presented by H-2Ld (P1CTL) (13) and have found that large unmodified tumors are highly resistant to therapy using the P1CTL (14, 15). In the process of studying the mechanism for resistance of large tumors, we have frequently observed recurrences of tumors in mice that have responded favorably to therapy with high numbers of transgenic T cells. To analyze the mechanism of tumor evasion of T cell therapy, we systematically characterized the P1A antigen among the recurrent tumors. We uncovered a large collection of tumor variants with mutations within the P1A epitope. These mutations abolished T cell recognition of the tumor cells by modulating either peptide binding to the MHC molecules or, more importantly, the binding of the MHC:peptide complex to the TCR. These results demonstrate antigenic drift of tumor antigens as a mechanism for tumor evasion of CTL therapy in vivo.

Methods
Experimental animals. Transgenic mice expressing a TCR specific for the tumor antigen H-2Ld:P1A35-43 complex have been described (13). TCR transgenic mice were backcrossed with BALB/cByj mice for at least nine generations before they were used for this study. BALB/c mice with a targeted mutation of the RAG-2 gene were purchased from Taconic Farms (Germantown, New York, USA).

Production of TCR α chain–transgenic mice. The transgenic vector consisting of the α chain of the TCR from a P1A-reactive CTL clone has been described (13). The non-TCR–related vector sequence was removed prior to injection into fertilized eggs of FVB/N mice. Founder mice were screened by PCR using primers for the rearranged VJ segment of the transgenic vector and by flow cytometry of the Vε8 chain on the surface of peripheral blood lymphocytes, as has been described (13).

Cell lines. H-2Ld–transfected, transporter associated with antigen processing 2–deficient RMA-S cells were produced and kindly provided by Ted Hansen (Washington University, St. Louis, Missouri, USA) (16). The BALB/c plasmocytoma J558 transfected with a pSV

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Nonstandard abbreviations used: T cell receptor (TCR); transgenic T cells expressing the α and β chains of TCR specific for tumor antigen P1A35-43 peptide presented by H-2Ld (P1CTL); PE, phycoerythrin.
vector (J558-Neo) has been described (14). To isolate tumor cells from ex vivo tissues, tumors were surgically removed and single-cell suspensions were prepared by grinding tumor tissues over two frosted glass slides. After removing the tumor debris, the viable cells were enriched by centrifugation through ficoll-paque medium. The tumor cells were cultured in RPMI medium containing 5% FCS for 1 week before they were used for flow cytometry and molecular analysis of the PIA gene.

Adoptive transfer of purified transgenic T cells. Pools of spleen and lymph node cells from P1CTL-transgenic mice were incubated with a cocktail of mAb’s (anti-CD4 mAb GK1.5, anti-FcR mAb 2.4G2, and anti-CD11c mAb N418). After removal of unbound mAb’s, the cells were incubated with anti-Ig–coated magnetic beads (BioSource International, Keystone, Colorado, USA). The Ab-coated cells were removed by a magnet. The unbound cells consisted of more than 90% CD8 T cells, with no detectable CD4 T cells. The purified CD8 T cells (5 × 10^6/mouse) were injected intravenously into mice bearing large tumors (>1 cm).

In other experiments, 2 × 10^7 spleen cells from either TCR α chain–transgenic mice (the F1 generation of a cross between transgenic founder and BALB/c mice) or their nontransgenic littermates were adoptively transferred into mice with tumors greater than 1 cm in diameter. Tumor growth was determined by physical examination, while the number of tumor-reactive T cells was monitored by flow cytometry.

Molecular characterization of PIA antigen from wild-type and CTL-resistant tumor cell lines. The expression of tumor antigen PIA was determined by RT-PCR using previously reported conditions (13). The primers used were 5′-GCTAGCTGGCGACCC-3′ (forward) and 5′-TTGGACCGTGAGCTTGAAGTGGAG-3′ (reverse). The PCR products were analyzed using agarose gel electrophoresis followed by cloning and sequencing.

To analyze the genetic lesion of the PIA antigen, the genomic DNA was isolated from either ex vivo tumor cells or their subclones, which were obtained by limiting dilutions. The PIA gene fragments were amplified by PCR using 5′-GCTAGCTGGCGACCC-3′ as the forward primer and 5′-GCTAGCTGGCGACCC-3′ as the reverse primer. The PCR products were cloned and sequenced. In some experiments, the PCR products were analyzed by digestion with a panel of restriction enzymes, namely AvaII, Fnu4HI, HaeIII, and BbsI, which distinguish wild-type and mutant PIA genes at positions 1, 6, 7, 8, and 9.

Peptide synthesis. The wild-type and mutant PIA peptides, as well as the control H-2L^d binding peptide from murine cytomegalovirus, were synthesized by Research Genetics (Huntsville, Alabama, USA), dissolved in ethanol, and stored at −20°C.

Stabilization of cell-surface H-2L^d on RMA-S–L^d. RMA-S–L^d cells were incubated with a given concentration of wild-type and mutant PIA peptides at 37°C for 16 hours in the presence of human β2-microglobulin (2.5 μg/ml; Sigma-Aldrich, St. Louis, Missouri, USA). Cell-surface H-2L^d was analyzed by flow cytometry.

Flow cytometry. Cell-surface expression of H-2L^d was detected using biotinylated mAb 28-14-8 (BD Pharmingen, San Diego, California, USA) followed by phycoerythrin-labeled (PE-labeled) streptavidin. To determine the binding of the H-2L^d–peptide complex to transgenic T cells, we used the H-2L^d-Ig dimer (purchased from BD Pharmingen) according to the manufacturer’s instructions. Briefly, 3 μg of peptides were incubated with 4 μg of H-2L^d-Ig and β2-microglobulin complex at 4°C for 48 hours in a total volume of 200 μl. PE-conjugated rat anti-mouse IgG1 mAb was added to the solution 1 hour before it was used to stain spleen cells from transgenic mice whose T cells expressed the TCR specific for the H-2L^d–PIA peptide complex. After washing away the unbound complex, the spleen cells were fixed with 1% paraformaldehyde in PBS and analyzed by flow cytometry.

Proliferation assay. To measure proliferation of transgenic T cells, spleen T cells (10^5/well) were cultured in 96-well plates in the presence of given concentrations of peptides. After 66 hours of culture, 1.25 μCi/well of 3H-thymidine was added. The cultures were harvested 12 hours later, and the incorporation of 3H-TdR was measured and used as an indicator of T cell proliferation.

Cytotoxicity assay. Transgenic spleen cells were stimulated with the P1A peptide (0.1 μg/ml) for 4 days and used as effectors. As targets, we used 51Cr-labeled P388D1 (H-2^d) cells. Various concentrations of peptides were added to the target cells immediately before the addition of the effector T cells. In the competition assay, the peptides added were a mixture of a given concentration of competing peptides and 10 ng/ml of the wild-type PIA peptide. The effector T cells and the targets were coincubated for 6 hours, and the percentages of specific lysis were calculated based on the following formula: specific lysis % = 100 × (cpm_sample – cpm_medium)/(cpm_max – cpm_medium).

Results

Monoclonal transgenic T cell therapy of large tumors in vivo selects for T cell–resistant tumor variants. It is generally accepted that large tumors are highly resistant to CTL therapy. To analyze the mechanism responsible for such resistance, we transplanted the plasmocytoma J558-Neo into syngeneic RAG-2−/−BALB/c mice. Once the tumors had reached a size of 1.2–1.4 cm in diameter, we adoptively transferred purified P1CTLs into tumor-bearing mice. At this point, the tumors would grow progressively in the untreated mice and euthanasia would be necessary within 1 week (14). Therapy with P1CTLs would prolong the life of the mice, although curing of large tumors was not achieved (14). As shown in Figure 1a, a reduction of more than 60% in tumor volume was observed within 1 week of T cell transfer. Histological examination revealed that the overwhelming majority of the tumor mass consisted
of necrotic tumor cells, although a small number of viable tumor cells was found (data not shown). However, tumor shrinkage stalled in the next 2 weeks. Starting from the third week, tumors resurfaced in all mice treated. We then started a new round of transfer of transgenic T cells. However, the tumors were no longer responsive to T cell treatment.

We isolated tumor cells from two of four mice that had received two rounds of CTL therapy and tested their susceptibility to cytolysis by activated transgenic T cells in vitro. The data in Figure 1b indicate that while parental tumor cells were readily killed by P1CTLs, those isolated from the two T cell–resistant tumors were completely refractory to treatment. RT-PCR analysis indicated that transcripts of tumor antigen P1A were expressed in both tumor lines (Figure 1c). Moreover, both lines expressed cell-surface H-2Ld at levels comparable to levels found on J558-Neo cells (Figure 1d). These results demonstrate that therapy with P1CTLs selects for tumor variants that are resistant to P1CTLs, and at least for the two resistant tumor lines tested here, resistance is not due to the lack of cell-surface MHC or loss of expression of the P1A gene.

Molecular lesions of P1A genes in tumor cells that evaded monoclonal T cell therapy. We isolated and sequenced five clones of P1A cDNA from each PCR product in Figure 1c. All cDNA clones from the J558-Neo tumors had the wild-type P1A sequence (Figure 2a), while those recovered from the CTL-resistant tumor cell lines harbored mutations within the P1A epitope, the nine-AA peptide corresponding to positions 35–43 of the predicted P1A protein (17). Four of five clones isolated from Tum 1 had a mutation that changed the AA sequence at position 6 (W→R, due to a point mutation of T→C) of the P1A epitope, herein termed P1A(6R). The other had a mutation that altered the AA sequence at position 9 (F→L, also due to a T→C point mutation), hereby termed P1A(9L). All five clones from Tum 2 were P1A(6R).

We carried out two types of experiments to identify the genetic lesions in the P1A gene in the two ex vivo tumor cell lines. First, using limiting dilutions, we obtained more than 30 clones from each of the parental J558-Neo, Tum 1, and Tum 2 cell lines, and then analyzed these for the presence of mutations at P6 and P9. Based on the mutations identified in the cDNA clones, we identified two restriction enzymes that can recognize the mutations. As indicated in Figure 2b, a mutation in the P6 position (T→C) rendered the P1A product susceptible to digestion by Fnu4HI, while a mutation in the P9 position (T→C) allowed recognition by AvaII. We therefore analyzed the genomic PCR products from multiple cell clones from parental J558-Neo, Tum 1, and Tum 2 cells for the presence of genetic lesions. As shown in Figure 2b, no PCR product from any of the 37 clones from J558-Neo cells harbored the P6 and P9 mutations. Further sequence analysis of multiple clones uncovered no mutations in the antigenic epitopes of the parental J558-Neo tumor cells, including those that were passed either in vitro or in vivo in untreated RAG-2–/– mice (data not shown). Thus, in the absence of a strong immune response, the P1A antigenic epitope remains unchanged.

The clones isolated from the Tum 1 line can be divided into four groups (b, lower panel). Group 1 (3/35) harbored a mutation at P9. Group 2 (3/35) appeared to be heterogeneous at position P9, while having no mutation at P6. Our analysis of the RNA transcripts derived from a representative clone from this group by RT-PCR revealed that the only form expressed was P1A(9L) (c). Thus the other potential allele was inactivated by some

![Figure 1](https://example.com/figure1.png)

**Figure 1** Adoptive therapy with P1A-specific transgenic T cells selects for T cell–resistant tumor cells. (a) Treatment of large P1A-expressing tumors with transgenic T cells results in rapid shrinkage of tumors, followed by stagnation and resurgence of tumor growth. J558-Neo tumor cells (5 × 10⁶) were injected subcutaneously into BALB/c mice. Two to three weeks later, when the tumors reached 1.2–1.4 cm in diameter, purified transgenic T cells (5 × 10⁶/mouse) were injected intravenously. The treatment was repeated on day 31 when the tumors had returned to their pretreatment size, as indicated by arrows. Mice were euthanized on day 39 (n = 4). (b) Tumor cells isolated after two treatments were resistant to cytolysis by activated P1A-specific T cells. Tumor cells were isolated from two different mice and were compared with their parental tumor J558-Neo cells for their susceptibility to cytolysis in a 6-hour cytotoxicity assay. Data shown are representative two experiments. A, AvaiI; F, Fnu4HI; U, uncut. (c) Expression of P1A was not lost in the P1CTL-resistant tumor cells. First-strand DNA was generated from total RNA using reverse transcriptase and amplified with primers specific for either GAPDH (700-bp product) or P1A (500-bp product). (d) Normal expression of H-2Ld on the surface of CTL-resistant tumor cells. Histograms depict binding of second-step reagent (dotted lines) or H-2Ld–specific mAb followed by the second-step reagent (solid lines). E/T, effector to target ratio.
yet-unidentified mechanism. Group 3 (27/35), which represented the majority of the clones isolated from Tum 1, had a mutation at P6. Group 4 members (2/35) are heterogeneous at P6 while having a wild-type sequence at P9. The cells in this group were lost prior to RNA analysis, and it is therefore unclear whether the apparent wild-type allele was also inactivated. Tum 2 clones can be divided into two groups, the majority of them (27/33) having a mutation at the P6 position, and the minority of them (6/33) harboring no mutation in either of these two positions (Figure 2b, upper panel). We also carried out extensive sequence analysis of the PCR products from Tum 1 and Tum 2. Our sequencing confirmed the mutations identified by enzymatic digestion. In addition, analysis of Tum 2 clones in which P1A mutations were not identified by enzymatic digestion revealed additional lesions in the antigenic epitope. Of the six clones from Tum 2 with no mutation at P6 and P9, one had a mutation at P7 (T→C, resulting in an L→P substitution), while five others harbored a mutation at P8 (T→G, leading to a V→G replacement), as shown in Figure 2a.

**Immunological basis of tumor evasion of monoclonal T cell therapy.** To test immunological consequences of these mutations, wild-type and mutant P1A peptides were synthesized and tested for their recognition by P1A-specific transgenic T cells. An H-2Ld binding peptide from murine cytomegalovirus was used as control (Figure 3a). As shown in Figure 3b, the mutant P1A(9L) peptide was 100-fold less potent than the wild type in inducing proliferation of transgenic T cells, while mutant P1A(7P) was about 1,000-fold less potent. Mutants P1A(6R) and P1A(8G) failed to induce any detectable T cell proliferation.

In a CTL assay to measure cytolysis of P388D1 target cells (Figure 3c), the P1A(9L) peptide was at least 10- to 100-fold less efficient than wild-type peptide, while P6 and P8 displayed a 10,000-fold reduction in inducing cytolysis of target cells. The most effective mutation was P1A(8G), which completely abolished recognition by P1CTLs.

Low-avidity TCR ligands can be TCR antagonists (18, 19). To test whether the mutant peptides are antagonists for P1A, we incubated varying concentrations of the mutant or wild-type P1A peptides, or unrelated H-2Ld-binding peptides, with the target cells in the presence of a fixed concentration of P1A peptide, and tested target-cell susceptibility to P1A-specific CTLs. As
shown in Figure 3d, P1A(9L) caused essentially no inhibition of P1A-mediated cytolysis. Moderate inhibition by P1A(6R), P1A(7P), and P1A(8G) was observed only when the mutant peptide was present at a 1,000-fold higher concentration than the P1A peptide. However, since the mutant peptides were significantly less potent than the unrelated peptide, which had no measurable avidity to TCR (see below), the observed inhibition by these mutant peptides was likely due to competition for H-2Ld binding rather than to its antagonism to TCR. Thus, these mutant peptides had no demonstrable antagonistic activity. Regardless of which mutations they harbored, all mutant J558-Neo clones were resistant to cytolysis by activated P1CTLs (Figure 4). Thus, under physiological conditions in which the antigenic peptides were produced at limiting doses, partial inactivation was sufficient to prevent T cell recognition of the tumor cells.

In theory, reduction of T cell recognition of the antigenic peptide can be attributed to either reduced peptide binding to MHC or reduced recognition of the MHC:peptide complex by the TCR. We determined the binding of wild-type and mutant peptides to H-2Ld using a classic H-2 stabilization assay with Ld-transfected RMA-S cells (16). The results are summarized in Figure 5. Interestingly, although P6 was not considered an anchor residue for H-2Ld-bound peptide, we found that P1A(6R) was somewhat less potent than the wild-type peptide in stabilizing H-2Ld. This is perhaps due to the positive charge introduced by the mutation. Similarly, replacing one permissible anchoring residue (F) with another (L) at P9 also had a moderate effect on peptide binding to H-2Ld. As expected, a control peptide from murine cytomegalovirus that had L at P9 reduced binding to H-2Ld. Surprisingly, mutations in P7 and P8 increased peptide binding to H-2Ld. All of the described alterations were statistically significant: P1A versus P1A(6R), \( P \leq 0.0001 \); P1A versus P1A(7P), \( P = 0.0492 \); P1A versus P1A(8G), \( P = 0.0316 \); P1A versus P1A(9L), \( P = 0.025 \).

To directly measure the effect of these mutations on T cell recognition of the H-2Ld:peptide complex, we loaded H-2Ldrg with a 166-fold excess of peptides and tested their binding to P1CTL. As shown in Figure 6a, in the spleen of the TCR transgenic mice, about 94–99% of CD8 T cells expressed high levels of the transgenic \( \alpha \) chain (no Ab against V\( \beta \)1 is available). About the same percentage of CD8 T cells were stained by the H-2Ld:P1A dimer. In contrast, only about 0.1% of CD8 T cells were stained by the control H-2Ld:peptide dimer. These results confirmed both the sensitivity and the specificity of the dimer binding assay. As shown in Figure 6b, both P1A(6R)- and P1A(9L)-loaded H-2Ld gave significant, albeit reduced, binding to the
transgenic T cells. A much-reduced binding was observed with H-2Ld:P1A(7P), while H-2Ld:P1A(8G) did not bind to P1CTLs. Titration of the MHC:peptide complex revealed that mutations P1A(6R) and P1A(9L) reduced the avidity between the TCR and the MHC peptide by about two- to fourfold; while H-2Ld:P1A(7P) showed a more than 30-fold reduction in binding to P1CTLs (Figure 6c). The reductions caused by all mutations were statistically significant (P1A versus P1A(6R), \( P = 0.035 \); P1A versus P1A(7P), \( P = 0.03 \); P1A versus P1A(8G), \( P = 0.0366 \); P1A versus P1A(9L), \( P = 0.0398 \)).

Polyclonal T cells with a transgenic TCR \( \alpha \) chain from a P1A-reactive CTL clone selected antigenic variants in vivo. To extend our observations from mice with essentially monoclonal T cells, we produced a transgenic mouse line with the TCR \( \alpha \) chain isolated from a P1A-reactive CTL clone. Since no transgenic \( \beta \) chain is present, the transgenic \( \alpha \) chain must pair with endogenous \( \beta \) chain to produce the cell-surface TCR/CD3 complex. As shown in Figure 7a, in the F1 generation of transgenic founder and BALB/c mice, more than 90% of the T cells expressed the transgenic TCR \( \alpha \) chain, as revealed by a V\( \alpha \)8-specific mAb. As a result, about 0.3% of CD8 T cells from TCR \( \alpha \) chain–transgenic mice bound to the H-2Ld:P1A complex (Figure 7a, lower panel), while no binding was seen in CD8 T cells from nontransgenic littermates (data not shown). To test whether T cells expressing the transgenic TCR \( \alpha \) chain can cause rejection of large tumors, we adoptively transferred spleen cells from TCR \( \alpha \) chain–transgenic mice into RAG-2–/– mice bearing large tumors. As shown in Figure 7b, the transgenic spleen cells, but not those from their nontransgenic littermates, caused rapid shrinkage of tumors. However, essentially all tumors recurred within a 4-week period. The tumor rejection response was associated with significant expansion of P1A-reactive CD8 T cells, as 10–70% of CD8 T cells in tumor-bearing mice were H-2Ld:P1A-specific (Figure 7c). We also analyzed the V\( \beta \) usage of the CD8 T cells that reacted to H-2Ld:P1A complex by flow cytometry using most of the anti-V\( \beta \) mAb's available. Our data demonstrate that, while the V\( \beta \) usage for the majority of the Ld:P1A-reactive T cells had not been identified due to limitation of mAb's, cells expressing V\( \beta \)3, V\( \beta \)4, V\( \beta \)6, V\( \beta \)8, V\( \beta \)10, V\( \beta \)11, V\( \beta \)12, and V\( \beta \)13 can be found among the Ld:P1A-reactive population (data not shown).

To determine whether antigenic variants can arise in mice with polyclonal P1A-reactive T cells, we isolated the genomic DNA from two recurrent tumors. The P1A gene fragments were amplified and then subjected to digestion by a panel of three restriction enzymes. Fnu4HI is specific to GCNGC and recognizes mutant P1A(6R). HaeIII, with a specificity for GGCC, can recognize mutations leading to L\( \rightarrow \)P replacements at either P7 or P1. Any change in the six nucleotides encoding P8 and P9 will result in loss of digestion by BbsI, which recognizes GTCTTC. As shown in Figure 8a, the P1A in tumors from nontransgenic T cell–treated mice was fully susceptible to BbsI while being resistant Fnu4HI and HaeIII. Thus, in the absence of antigen-specific T cells, no mutations in the P1A epitope could be identified by restriction enzyme mapping. In contrast, the tumor cells recovered from mice treated with TCR \( \alpha \) chain–transgenic T cells yielded P1A products that were partially resistant to BbsI and partially susceptible to Fnu4HI. Thus, a substantial number of tumor cells harbored the mutation P1A(6R). To determine the mutations that rendered resistance to BbsI, we cloned the PCR products and sequenced three of the clones that were resistant to BbsI. Two clones had a
change of T→G, resulting in the P1A(8G) mutant, while the other had a mutation of G→C, which leads to a substitution of V with L at P8, hereby called P8L. Since mutations at the P8 and P6 positions abolished T cell recognition of T cells, we conclude that antigenic drift also allows tumor evasion of polyclonal CTL responses, predominantly against the P1A epitope.

Discussion
Although antigenic drift has been established as a major mechanism for viral evasion of host immunity (1–5), its contribution to tumor evasion has not been systematically analyzed. There are at least three constraints that may have hindered progress in this area. First, given the rapid replication and — in the cases of RNA viruses and retroviruses — instability of the viral genome, an infected host can conceivably have a large number of mutant viral genomes available for immune selection. Second, perhaps because viruses are foreign to the host, immune responses are generally more robust than antitumor immunity, in terms of both the number of virus-specific CTLs (20–22) and their effector function (13, 23). Third, in no experimental tumor model has the spectrum of tumor rejection antigens been fully characterized, despite the fact that many tumor antigens have been identified (24). The lack of clarity in the spectrum of tumor-rejection antigens makes it difficult to analyze mutations that may allow tumor evasion.

In order to separate these three intertwining issues, we used transgenic mice with either TCR α or TCR αβ transgenes to determine whether tumors are genetically versatile enough to evade CTLs specific for a natural tumor antigenic epitope. Using T cells from mice with monoclonal (αβ chain–transgenic) and polyclonal (α chain–transgenic) P1A-reactive T cells, we obtained clear evidence that mutations in at least four residues within a nine-AA peptide can each result in tumor evasion of a robust CTL response in vivo, as tumors with a single mutation were shown to recur and progressively grow in the presence of a strong CTL response.

It is of interest to note that of almost 70 clones of ex vivo tumor variants analyzed, apparent heterozygosity is found in only five clones. Since mutation in both alleles in the majority of the clones is unlikely, we believe the J558-Neo tumor cells probably have only one copy of the P1A gene to begin with. Since P1A was recently mapped to the X chromosome (25), the hemizygosity of the gene may be due to the fact that the tumor cells are
derived from male mice (although our PCR analysis for a Y-chromosomal marker indicated that J558-Neo cells lack the Y chromosome, data not shown) or to the loss of heterozygosity of five clones during tumorigenesis. The apparent heterozygosity may be due to the accidental seeding of two independent clones into one well.

A related issue is whether the J558-Neo cells used for the current study are comprised of a mixture of cells with a high frequency of mutation in P1A antigenic epitopes. We believe this is unlikely for three reasons. First, our analysis of 37 clones from parental cells by restriction enzyme mapping and sequencing revealed no mutations in the antigenic epitope. Second, our repeated efforts to select for P1A mutants with activated P1A-specific CTLs and parental tumor cells in ten 96-well plates (5 × 10^4 tumor cells/well) have failed to obtain any CTL-resistant clones in vitro (data not shown). This may put the frequency of mutants at less than one per 5 × 10^7 cells. These results do suggest that cells with mutations in the P1A epitope are not present at high frequencies prior to T cell selection. Since we started T cell therapy only after tumors had reached a size of more than 1.2 cm in diameter, we estimate that there may be up to 10^9 tumor cells per mouse at the time of treatment. As a result, there could be a significant number of tumor variants for in vivo selection. Third, tumor cells isolated from RAG-2^{-/-} mice that received no T cell therapy did not reveal mutations in the P1A antigenic epitope (data not shown). Likewise, tumor cells recovered from mice that received nontransgenic T cells also contained no mutations in the P1A epitope. Therefore, the mutants described here must result from immune selection.

Our work demonstrated that, in addition to antigen loss (lack of expression of genes encoding tumor antigen) (12, 26) and MHC loss (10, 11), the antigenicity of tumors can be altered by antigenic drift. Thus, much like viruses, tumor cells can evade T cell–mediated destruction through mutations in their antigenic epitopes. Although we are not aware of direct evidence, two lines of previous work are consistent with the notion that mutations in tumor antigenic epitopes can be a mechanism for immune evasion by tumor cells. First, CTL clones have long been used as tools to isolate tumor antigenic mutants in vitro, although long and repeated selection in vitro is usually needed for this purpose (17, 27, 28). Second,
of the antigenic loss variants reported in the classic study (12), one was later proven to contain a mutation in the P1A epitope (position 7, V→A) (17). This mutation prevented the clone from being recognized by one, but not another, CTL clone specific for the same epitope (17). However, the authors reported that the tumor clone harboring this mutation had a significant growth advantage in vitro over other tumor cells in the same isolate (12). It is thus unclear whether this tumor variant was selected for in vivo, and if so, whether it was selected based on its lack of antigenicity. If both these questions were answered in the affirmative, it would indicate that antigenic drift in tumors can occur in mice with an unmanipulated T cell repertoire.

Since the majority of our data involves adoptive transfer of purified transgenic T cells into immune-deficient mice, it is of interest to consider whether antigenic drift can occur in a normal tumor-bearing host in which the immune response consists of CD4 T cells, CD8 T cells, and B cells. Since we have verified our conclusion with adoptively transferred total spleen cells from TCR α chain–transgenic mice that have all subsets of lymphocytes, we believe antigenic drift can happen in a host with multiple subsets of lymphocytes. Moreover, recent studies indicate that homeostatic proliferation of T cells, upon adoptive transfer to an immune-deficient host, may enhance the efficacy of immune therapy both in experimental animals (29) and in humans (30). Such a setting is essentially identical to the approach in the current study.

Several properties of the P1A antigenic variants deserve comment. First, although there is a general correlation between the binding of the H-2Ld:peptide complex to the TCR and the ability of the peptides to induce T cell responses, the difference in binding is considerably less than that in biological function. For instance, mutation at P6 reduces binding by less than fourfold, yet it reduces the activity of the peptide in inducing proliferation and cytolysis by more than 10,000-fold. The fact that a moderate difference in avidity of a TCR-ligand interaction has a drastic effect on the consequence of T cell recognition has been described and discussed extensively (31). Second, and more surprisingly, the intensity of MHC:peptide binding to a TCR does not always correlate well with biological function of the peptide. Thus H-2Ld:P1A(7P) binds much less efficiently than H-2Ld:P1A(6R) does, yet it is P1A(7P), not P1A(6R), that can induce T cell proliferation. One possible explanation is the higher potency of P1A(7P) in binding H-2Ld. Likewise, although P1A(6R) and P1A(9L) have comparable binding to H-2Ld and TCR, there was at least a 100-fold difference in biological activity between the two mutants. Davis et al. suggested that induced aggregation of MHC complex might be involved in T cell activation (32). In this scenario, different MHC:peptide complexes may differ in their ability to aggregate after TCR engagement, and the extent of such aggregation may determine the consequences of interaction between TCR and MHC:peptide. Finally, although some mutations do not completely abolish T cell recognition of the P1A peptide, cells harboring such mutations are completely resistant to CTL lysis. This is most likely due to limiting amounts of peptides produced intracellularly and explains the ability of these tumor mutants to evade T cell recognition.

Despite our extensive efforts, we have not been able to identify antigenic variants in the P1A gene in recurrent tumors isolated from mice that received therapy with primed nontransgenic T cells (data not shown). However, we believe that this is due to our incomplete understanding of the nature of the T cells responsible for tumor rejection. Better characterization of the antigenic spectrum will also allow determination of whether antigenic drift can lead to epitope spreading. Nevertheless, the results reported here reveal that T cell therapy directed against a known antigenic epitope can be rendered ineffective by selection of tumor antigenic variants regardless of whether the T cells are polyclonal or monoclonal. These results demonstrate that, much like viruses, tumor cells can evade T cell–mediated destruction through mutation in their antigenic epitope. Given the fact that a similar finding over 10 year ago (3) with transgenic T cells specific for one viral epitope has now been extended into a variety of in vivo models (4, 33, 34), further characterization
of tumor antigens should allow testing of whether the “antigenic drift” of tumor cells described here can be generalized to cancer patients. In this regard, it is worth noting that recent studies revealed the coexistence of tumor-reactive T cells and tumors in cancer patients (21, 22), even in the draining lymph nodes (35). In many cases, the mRNA encoding tumor antigens was retained (35). It would be of interest to analyze whether the antigenic epitopes were mutated in these cancer patients. Regardless of whether tumor cells can evade T cell responses to more than one epitope, the fact that mutations in a single epitope can occur suggests that vaccination based on a limited set of peptides (36) and gene therapy using TCR with monospecificity may have limited value (37).

Surprisingly, a recent study demonstrated that a mostly monoclonal T cell response develops over time, even if the initial T cells are polyclonal in nature (30). This may be overcome either by immunization with a multitude of tumor antigens (38–40) or by choosing those antigens that must be retained if the cancer cells are to remain cancerous (41).

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