Adoptive T Cell Therapy for Cancer: Boutique Therapy or Treatment Modality?

Running Title: TIL Therapy as a treatment modality

Cassian Yee MD  
Professor, Dept of Melanoma Medical Oncology  
Professor, Immunology  
UT MD Anderson Cancer Center  
7455 Fannin St, Unit #904  
Houston TX  
77054

Contact email:  
cyee@mdanderson.org  
713 563 3750

Conflict of Interest Statement:  
The author has no conflicts relevant to the content of this manuscript

Summary
Adoptive cellular therapy, involving the ex vivo enrichment and expansion of antigen-specific immune cells for adoptive transfer, has emerged as an increasingly effective modality for the treatment of patients with advanced cancer refractory to conventional therapy.

In this issue of CLINICAL CANCER RESEARCH, Besser and colleagues present an intent-to-treat (ITT) analysis of 80 patients enrolled on an adoptive therapy study using ex vivo expanded unselected tumor-infiltrating lymphocytes (1). Among this cohort of patients with advanced disease, the majority presenting with poor prognostic metastatic melanoma (Stage M1c), 57 were eventually treated, with about half of those not receiving treatment due to non-clinical reasons (no TIL growth or refusal to further participate) and half clinically progressing too rapidly to receive therapy. Overall response rate and median survival was about 40% and 15 months among the 57 treated patients and 30% and 10 months for all enrolled. Considering that all patients had had at least one previous treatment for metastatic disease, (often multiple prior lines of aggressive therapy), and the natural history of melanoma affecting visceral sites, these are very encouraging results for patients and for the field of immune-based therapies in general.

The development of immunotherapies for the treatment of refractory or recurrent disease has witnessed a renaissance of late in both cell-based and immunomodulatory approaches. Clinical trials using antibodies to establish immune checkpoint blockade against CTLA4 and the PD-1/PDL-1 axis report significant longlasting responses via in vivo activation and expansion of the endogenous anti-tumor immune response (2). As a means of providing an exogenous source of ex vivo expanded effector cell, adoptive
cellular therapy, has also emerged as a highly effective modality capable of eliciting durable and complete responses.

Three forms of adoptive cellular therapy using T cells have been practiced—TIL therapy, using lymphocytes expanded from a tumor biopsy sample (3), antigen-specific T cell therapy, using endogenous T cells sourced from peripheral blood (4-6), and more recently, the use of gene-modified T cells engineered to express the desired TCR or chimeric antigen receptor (CAR) with occasional remarkable results (7) (Figure 1).

When it became apparent that the duration of T cell survival after adoptive transfer correlated with clinical response; strategies to enhance in vivo persistence were implemented involving both extrinsic modification of the host environment, through the use of conditioning lymphodepletion or intrinsic manipulation of the effector T cell itself, by enhancing cellular replicative potential via cytokine modulation (8), phenotype-based selection or genetic engineering (7). By incorporating these approaches into the TIL therapy protocol, a significant increase in clinical response rates was achieved (> 50% in select cases) and durable complete remissions in the setting of significant tumor burden (9).

In the field of adoptive T cell therapy using TIL, two important milestones were attained, in large part through pioneering efforts of the Surgery Branch at the NCI, enabling its promulgation into the clinical arena as a feasible therapeutic option: one was a means of expanding the TIL population 1000-5000 fold based on methods originally established for antigen-specific T cell expansion using a TCR trigger (anti-CD3) and irradiated feeder cells, and the second, was inclusion of a lymphodepleting conditioning regimen for patients prior to TIL infusion. This regimen was initially nonmyeloablative, and later advanced to a TBI-containing ablative regimen with a commensurate increase in serious adverse toxicities, but also dramatic and durable clinical responses (up to 40%). The nonmyeloablative regimen used in this study is the most established; coupled with an expedited protocol to generate ‘young’ TIL that was successfully expanded and infused in more than 90% of patients, this represented a potential ‘standardized’ protocol with which to move forward to a randomized controlled clinical trial given the encouraging ITT results.

The impetus to develop improved and simpler TIL protocols arose from prior work at the NCI and other centers involved in larger scale TIL trials such as those conducted at MD Anderson Cancer Center (10) and at Sheba Medical Center (1) where response rates of 40% or more were consistently achieved among patients who eventually received treatment. Although these studies corroborated the original promising results, only 40% to as few as 27% of patients who underwent resection for TIL generation ultimately received TIL therapy (11) this attrition due in part to disease progression, protocol-specific and product-related exclusion criteria - features which could be addressed by a shortened time to therapy from enrollment and modification of product release criteria. While the original TIL protocol, commonly practiced at the NCI required 7-8 weeks from resection to TIL product, the young TIL protocol developed by Tran et al shortened the pre-expansion phase, eliminated exclusion of TIL cultures based on absent in vitro activity and produced a TIL product in 4 weeks (12); implementation of this protocol in the study presented by Besser et al here, led to > 70% treatment: enrollment ratio, with response rates among treated patients at least as favorable as those
shown in prior studies. Only 10% of patients failed to yield a useable TIL product and 14% were excluded due to clinical deterioration.

There remain however a number of issues yet to be resolved: should ablative radiation therapy be added to the conditioning regimen and which patients should be considered for this life-threatening but highly effective treatment? Can a superior TIL effector population be defined on the basis of in vitro phenotype selection or cytokine modulation? Is there a clinical or immunological biomarker profile that can identify patients predicted to respond to therapy? Furthermore, the treatment landscape for patients with metastatic melanoma has changed in a very positive and dramatic fashion over the last 5 years. With the advent of more and more positive data arising from the use of immune checkpoint inhibitors and targeted therapies, alone, in combination with each other, or in combination with conventional modalities, it is becoming less and less obvious which algorithmic endpoint cellular therapies will eventually find its niche; more than likely, combinational therapies involving the use of clinically-approved immunomodulators together with adoptive cellular therapies will be established as the standard of practice for clinical trials and hopefully, a standard of care.

In the parlance, then, of today’s Youtube generation, is adoptive cellular therapy ready to go ‘viral’ as in the case of a recently popularized Korean music video star, or is it limited still to an eclectic group of diehard believers? The answer lies somewhere in between. While there remains much to be addressed by taking a reductionist approach to adoptive cellular therapy - by isolating and expanding a uniform population of antigen-specific T cells, epigenetically modulating or genetically engineering an ideal central memory / stem cell effector population, limiting toxicities and fine tuning affinities - there is reason to believe that cellular therapy is now poised to make the leap from ‘boutique therapy’ to ‘treatment modality’. The report presented here describes one significant step towards this goal-and now, may just be the time for adoptive cellular therapy to go mainstream.

**Figure Legend**

**Figure 1. Adoptive Cell Therapy**

Adoptive Cell Therapy is represented by three general approaches:

1) Enrichment and expansion of **tumor-infiltrating lymphocytes (TIL)** from a disaggregated tumor biopsy sample

2) Genetic transfer of **T Cell Receptor (TCR)** recognizing tumor antigen-derived peptide-MHC target or **Chimeric Antibody Receptor (CAR)** recognizing surface tumor protein

3) Enrichment of endogenous **antigen-specific T cells** from peripheral blood mononuclear cells by in vitro stimulation followed by cell selection or cloning. PBMCs are a source of both antigen-presenting cells and T cells.

Following enrichment, the population of tumor-reactive T cells undergoes rapid expansion of 1000-5000 fold achieving 10-100 billion cells for adoptive transfer. Patients often receive a lymphodepleting conditioning regimen pre-infusion followed by exogenous IL-2. In the case of adoptive TIL therapy, patients receive high-dose near ablative or fully ablative conditioning pre-infusion and a course of high-dose IL-2 post-infusion. In Besser et al, ‘young’ TIL are generated using a shortened pre-expansion
culture phase prior to rapid cell expansion, enabling production of an infusible T cell product within 5-7 weeks from time of tumor collection.
References


Figure 1:

TIL

Treatment modality
Source of T cells

TCR/CAR

Ag-presenting cells

T cell

Antigen-specific T-cell enrichment

PBMC

Cloning, selection or sort

TCR + zeta

Chimeric Ab + zeta

Transfection of PBMC

Conditioning

T-cell expansion and infusion
Adoptive T Cell Therapy for Cancer: Boutique Therapy or Treatment Modality?

Cassian Yee

Clin Cancer Res Published OnlineFirst August 6, 2013.

Updated version
Access the most recent version of this article at:
doi:10.1158/1078-0432.CCR-13-1367

Supplementary Material
Access the most recent supplemental material at:
http://clincancerres.aacrjournals.org/content/suppl/2013/09/04/1078-0432.CCR-13-1367.DC1

Author Manuscript
Author manuscripts have been peer reviewed and accepted for publication but have not yet been edited.

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.