Clinical responses with T lymphocytes targeting malignancy-associated κ light chains

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**BACKGROUND.** Treatment of B cell malignancies with adoptive transfer of T cells with a CD19-specific chimeric antigen receptor (CAR) shows remarkable clinical efficacy. However, long-term persistence of T cells targeting CD19, a pan–B cell marker, also depletes normal B cells and causes severe hypogammaglobulinemia. Here, we developed a strategy to target B cell malignancies more selectively by taking advantage of B cell light Ig chain restriction. We generated a CAR that is specific for the κ light chain (κ.CAR) and therefore recognizes κ-restricted cells and spares the normal B cells expressing the nontargeted λ light chain, thus potentially minimizing humoral immunity impairment.

**METHODS.** We conducted a phase 1 clinical trial and treated 16 patients with relapsed or refractory κ+ non-Hodgkin lymphoma/chronic lymphocytic leukemia (NHL/CLL) or multiple myeloma (MM) with autologous T cells genetically modified to express κ.CAR (κ.CARTs). Other treatments were discontinued in 11 of the 16 patients at least 4 weeks prior to T cell infusion. Six patients without lymphopenia received 12.5 mg/kg cyclophosphamide 4 days before κ.CART infusion (0.2 × 10⁸ to 2 × 10⁸ κ.CARTs/m²). No other lymphodepletion was used.

**RESULTS.** κ.CART expansion peaked 1–2 weeks after […]

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Clinical responses with T lymphocytes targeting malignancy-associated κ light chains

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BACKGROUND. Treatment of B cell malignancies with adoptive transfer of T cells with a CD19-specific chimeric antigen receptor (CAR) shows remarkable clinical efficacy. However, long-term persistence of T cells targeting CD19, a pan–B cell marker, also depletes normal B cells and causes severe hypogammaglobulinemia. Here, we developed a strategy to target B cell malignancies more selectively by taking advantage of B cell light Ig chain restriction. We generated a CAR that is specific for the κ light chain (κ.CAR) and therefore recognizes κ-restricted cells and spares the normal B cells expressing the nontargeted λ light chain, thus potentially minimizing humoral immunity impairment.

METHODS. We conducted a phase 1 clinical trial and treated 16 patients with relapsed or refractory κ- or non-Hodgkin lymphoma/chronic lymphocytic leukemia (NHL/CLL) or multiple myeloma (MM) with autologous T cells genetically modified to express κ.CAR (κ.CARTs). Other treatments were discontinued in 11 of the 16 patients at least 4 weeks prior to T cell infusion. Six patients without lymphopenia received 12.5 mg/kg cyclophosphamide 4 days before κ.CART infusion (0.2 × 10^8 to 2 × 10^8 κ.CARTs/m²). No other lymphodepletion was used.

RESULTS. κ.CART expansion peaked 1–2 weeks after infusion, and cells remained detectable for more than 6 weeks. Of 9 patients with relapsed NHL or CLL, 2 entered complete remission after 2 and 3 infusions of κ.CARTs, and 1 had a partial response. Of 7 patients with MM, 4 had stable disease lasting 2–17 months. No toxicities attributable to κ.CARTs were observed.

CONCLUSION. κ.CART infusion is feasible and safe and can lead to complete clinical responses.

TRIAL REGISTRATION. ClinicalTrials.gov NCT00881920.

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Introduction

Chimeric antigen receptors (CARs) most commonly combine the antigen-recognizing portion of a mAb with the signaling machinery of T cells. When these molecules are genetically grafted into T lymphocytes, they allow these cells to bind to tumor surface antigens in an MHC-independent fashion and promote T cell costimulation, thereby activating their cytotoxic pathways against the malignant cells (1). Autologous T cells expressing CD19-directed CARs (CD19.CARTs) have shown remarkable activity in patients with acute lymphoblastic leukemia (ALL) (2–6) and some activity in patients with chronic lymphocytic leukemia or small lymphocytic lymphoma (CLL/SLL) (7–9) or non-Hodgkin lymphoma (NHL) (10–13). Nonetheless, sustained clinical responses due to CD19.CARTs usually require long-term persistence and invariably are associated with B cell aplasia; CD19 is a pan–B cell marker, and normal B lymphocytes are also a target for CD19.CART cytotoxicity (5).

If it were possible to effectively target antigens that have a more restricted distribution on malignant B cells versus normal B lymphocytes, then sufficient numbers of normal B lymphocytes might be spared to avoid profound hypogammaglobulinemia. We therefore took advantage of the surface expression of Igks on mature B lymphocytes and mature B lymphoid malignancies, since each cell expresses either κ or λ light chains, but not both. We reasoned that targeting the light chain subtype expressed by monoclonal malignant NHL and CLL/SLL B cells should efficiently kill tumor cells, while sparing normal B cells expressing the reciprocal subtype. This concept is also potentially applicable to multiple myeloma (MM). Although plasma cells no longer express surface Igks, several groups have described an MM-initiating population that expresses surface Igks (14). We have previously devel-
developed a CAR that targets the κ light chain (κ.CAR) (15), and here we show results of a phase 1 clinical trial using autologous T cells genetically modified to express the κ.CAR (κ.CARTs). We demonstrate that this approach is feasible and safe and can lead to complete clinical responses.

**Results**

*Patients' characteristics.* We administered κ.CARTs to 16 patients, 9 with NHL (including CLL/SLL) and 7 with MM (Figure 1). Their characteristics are summarized in Tables 1 and 2. All patients had active disease at the time of κ.CART infusion, and all had relapsed after one or more lines of chemotherapy. Six patients who had neither received other chemotherapy within 4 weeks of κ.CART infusion nor had absolute lymphocyte counts (ALCs) above 500/μl received 12.5 mg/kg cyclophosphamide (CTX) 4 days before the first infusion of κ.CARTs. Five patients received 1 final cycle of chemotherapy within 4 weeks of κ.CART infusion, as documented in Tables 1 and 2. These chemotherapeutic regimens had been previously chosen by the referring physicians as salvage treatment and were not specifically selected for their lymphodepleting potential.

**Characteristics of infused CARTs.** We manufactured 27 κ.CART lines from peripheral blood mononuclear cells (PBMCs) isolated from 24 patients (Figure 1). One patient had 2 cell lines and another patient had 3 cell lines made to allow repeat infusions. CAR transduction levels were adequate (>80%) in all manufactured T cell products, regardless of the cytokine(s) used for the expansion (Supplemental Figure 1B; supplemental material available online with this article; doi:10.1172/JCI86000DS1). All lines were composed of more than 99% CD3+ T cells, and each contained a variable ratio of CD4+ and CD8+ cells, with an overall CD8 predominance in the T cell products expanded in IL-2 (78% ± 11%) compared with those expanded in IL-7 and IL-15 (56% ± 12%; P < 0.001) (see Methods and Supplemental Figure 1C). The majority of κ.CARTs were CD45RO+ and lacked CCR7, but a small fraction expressed the central memory–associated phenotypic markers CD62L, CD27, and CD28 (Supplemental Table 1). T cell products grown with IL-7 and IL-15 had approximately twice as many CD45RA+ cells as those grown in IL-2 (4.9% ± 4.1% vs. 10.7% ± 7.1%, P = 0.03) (Supplemental Table 1). NK cells (CD3 CD56+) were not detectable. All products demonstrated cytotoxicity against κ+ targets in vitro, as assessed by 51Cr-release assays (Supplemental Figure 1D). Cytotoxic activity against κ targets was negligible. Eight patients did not receive their cell lines (Figure 1), because they pursued alternative treatment outside our institution (6 patients), had progressive disease (PD) during cell manufacture (1 patient), or had no evidence of disease at the time of planned infusion (1 patient).

**κ.CART expansion and persistence.** Molecular signals for κ.CARTs were consistently detected in the peripheral blood at low levels 3 hours after the first κ.CART infusion but increased to peak at 1 to 2 weeks after the first infusion (94 ± 186 copies/μg of PBMC DNA and 248 ± 750 copies/μg of DNA at weeks 1 and 2, respectively, for all patients). The signals then declined to lower levels by week 6 (26 ± 49 copies/μg of DNA) (Figure 2). Seven patients received a second infusion. The difference in expansion between the first and second infusions was not statistically significant (AUCs for 2 and 6 weeks after infusion: P = 0.109 and P = 0.219, Wilcoxon signed-rank test). Transgene levels were always below the threshold needed to detect a distinct CART population by flow cytometry and became almost undetectable 6 months after the last infusion (3 ± 4 copies/μg of PBMC DNA). There was no correlation between transgene levels in the peripheral blood and preinfusion ALC, disease type, cell dose, cytokines used in culture (IL-2 vs. IL-7 and IL-15), or clinical response (data not shown).
hypogammaglobulinemia as a result of prior treatment, the biology of their underlying disease, or both. With the exception of the 2 CLL patients, B cell lymphopenia was especially pronounced in the NHL patient group (with 1.5% ± 2% of CD19+ cells gated on CD45+ lymphocytes at the time of infusion). This made assessment of new-onset B lymphopenia due to κ.CARTs challenging.

Acute and long-term toxicities. All infusions were well tolerated. Apart from a patient with MM who had a grade 3 lymphopenia that was deemed possibly related, none of the adverse events reported (most frequently anemia, leukopenia, fatigue, hyper- or hypokalemia, and elevated aspartate aminotransferase [AST]) were considered to be related to κ.CART infusion. The vast majority of patients had baseline B cell lymphopenia and polyclonal hypogammaglobulinemia as a result of prior treatment, the biology of their underlying disease, or both. With the exception of the 2 CLL patients, B cell lymphopenia was especially pronounced in the NHL patient group (with 1.5% ± 2% of CD19+ cells gated on CD45+ lymphocytes at the time of infusion). This made assessment of new-onset B lymphopenia due to κ.CARTs challenging. However, in patients with MM, who consistently had more

### Table 1. Characteristics of patients with NHL or CLL

<table>
<thead>
<tr>
<th>UPIN</th>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Previous therapies</th>
<th>Cytokines in CART culture</th>
<th>Time from last chemo. treatment</th>
<th>Pre-CART CTX</th>
<th>DL</th>
<th>CAR+ cells in product (%)</th>
<th>CAR+ T cells/m2 admin.</th>
<th>No. of infusions</th>
<th>Best response</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>53</td>
<td>F</td>
<td>LPL</td>
<td>R-CHOP, 2CDA, R-Beam/ASCt, dexamethasone/bortezomib</td>
<td>IL-2</td>
<td>9 wk</td>
<td>Yes</td>
<td>1</td>
<td>87</td>
<td>1.7 × 107</td>
<td>1</td>
<td>NR</td>
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<tr>
<td>2</td>
<td>59</td>
<td>M</td>
<td>FL/DLBCL</td>
<td>R-CHOP/XRT, FCR, R-IEE, TTR, CD95/CART, R-bendamustine</td>
<td>IL-2</td>
<td>4 wk</td>
<td>No</td>
<td>1</td>
<td>85</td>
<td>1.7 × 107</td>
<td>3</td>
<td>CR × 32+ mo</td>
</tr>
<tr>
<td>3</td>
<td>70</td>
<td>M</td>
<td>DLBCL</td>
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<td>IL-2</td>
<td>6 d</td>
<td>No</td>
<td>2</td>
<td>90</td>
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<td>1</td>
<td>NR</td>
</tr>
<tr>
<td>4</td>
<td>72</td>
<td>M</td>
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<td>R-bendamustine</td>
<td>IL-2</td>
<td>4 wk</td>
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<td>2</td>
<td>91</td>
<td>9.1 × 107</td>
<td>1</td>
<td>NR</td>
</tr>
<tr>
<td>5</td>
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<td>M</td>
<td>LPL</td>
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<td>IL-2</td>
<td>1 yr</td>
<td>No</td>
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<td>94</td>
<td>1.9 × 107</td>
<td>2</td>
<td>PR × 3 mo</td>
</tr>
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<td>55</td>
<td>M</td>
<td>FL/DLBCL</td>
<td>R-CHOP, R-IEE, R-Beam/ASCt, rituximab</td>
<td>IL-7/IL-15</td>
<td>16 wk</td>
<td>No</td>
<td>3</td>
<td>85</td>
<td>1.7 × 107</td>
<td>6</td>
<td>CR × 6 wk</td>
</tr>
<tr>
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<td>CLL/SLL</td>
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<td>IL-7/IL-15</td>
<td>1 yr</td>
<td>No</td>
<td>3</td>
<td>86</td>
<td>1.6 × 107</td>
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<td>SD × 6 wk</td>
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<td>8</td>
<td>75</td>
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<td>MCL</td>
<td>R-NCVAQD, bortezomib, carfilzomb/lenalidomide, R-bendamustine</td>
<td>IL-7/IL-15</td>
<td>20 d</td>
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<td>3</td>
<td>86</td>
<td>1.8 × 107</td>
<td>1</td>
<td>NR</td>
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<tr>
<td>9</td>
<td>69</td>
<td>M</td>
<td>DLBCL</td>
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<td>14 d</td>
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<td>3</td>
<td>93</td>
<td>1.9 × 107</td>
<td>1</td>
<td>NR</td>
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<tr>
<td>10</td>
<td>69</td>
<td>F</td>
<td>CLL/SLL</td>
<td>R-fludarabine, R-bendamustine</td>
<td>IL-7/IL-15</td>
<td>1 yr</td>
<td>No</td>
<td>3</td>
<td>86</td>
<td>1.6 × 107</td>
<td>2</td>
<td>SD × 6 wk</td>
</tr>
<tr>
<td>11</td>
<td>54</td>
<td>M</td>
<td>κVAD</td>
<td>melphalan/ASCT</td>
<td>IL-2</td>
<td>34 wk</td>
<td>Yes</td>
<td>2</td>
<td>92</td>
<td>9.2 × 107</td>
<td>1</td>
<td>SD × 17 mo</td>
</tr>
<tr>
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<td>60</td>
<td>M</td>
<td>κVAD</td>
<td>melphalan/ASCT</td>
<td>IL-2</td>
<td>27 wk</td>
<td>Yes</td>
<td>3</td>
<td>76</td>
<td>1.5 × 107</td>
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<td>SD × 24 mo</td>
</tr>
<tr>
<td>13</td>
<td>58</td>
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<td>κVAD</td>
<td>melphalan/ASCT</td>
<td>IL-2</td>
<td>2.5 yr</td>
<td>Yes</td>
<td>3</td>
<td>92</td>
<td>1.8 × 104</td>
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<td>SD × 6 wk</td>
</tr>
<tr>
<td>14</td>
<td>54</td>
<td>M</td>
<td>κVAD</td>
<td>melphalan/ASCT, VDT-PC, VDT-AC</td>
<td>IL-7/IL-15</td>
<td>5 d</td>
<td>No</td>
<td>3</td>
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<td>69</td>
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<td>IL-7/IL-15</td>
<td>15 wk</td>
<td>Yes</td>
<td>3</td>
<td>90</td>
<td>1.9 × 104</td>
<td>2</td>
<td>SD × 6 wk</td>
</tr>
<tr>
<td>16</td>
<td>43</td>
<td>F</td>
<td>κVAD</td>
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<td>IL-7/IL-15</td>
<td>19 d</td>
<td>No</td>
<td>3</td>
<td>93</td>
<td>1.9 × 104</td>
<td>1</td>
<td>NR</td>
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<tr>
<td>17</td>
<td>59</td>
<td>F</td>
<td>κVAD</td>
<td>melphalan/ASCT, R-Beam/ASCt, dexamethasone/bortezomib</td>
<td>IL-7/IL-15</td>
<td>28 wk</td>
<td>Yes</td>
<td>3</td>
<td>94</td>
<td>1.8 × 104</td>
<td>2</td>
<td>SD × 6 wk</td>
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</tbody>
</table>

### Table 2. Characteristics of patients with MM

<table>
<thead>
<tr>
<th>UPIN</th>
<th>Age</th>
<th>Sex</th>
<th>Myeloma subtype</th>
<th>Previous therapies</th>
<th>Cytokines in CART culture</th>
<th>Time from last chemo. treatment</th>
<th>Pre-CART CTX</th>
<th>DL</th>
<th>CAR+ cells in product (%)</th>
<th>CAR+ T cells/m2 admin.</th>
<th>No. of infusions</th>
<th>Best response</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>56</td>
<td>M</td>
<td>IgGκ</td>
<td>RVD, melphalan/ASCT</td>
<td>IL-2</td>
<td>34 wk</td>
<td>Yes</td>
<td>2</td>
<td>92</td>
<td>9.2 × 107</td>
<td>1</td>
<td>SD × 17 mo</td>
</tr>
<tr>
<td>7</td>
<td>53</td>
<td>F</td>
<td>IgGκ</td>
<td>RD, VD, RVD, CyBorD, Melphalan/ASCT</td>
<td>IL-2</td>
<td>27 wk</td>
<td>Yes</td>
<td>3</td>
<td>76</td>
<td>1.5 × 107</td>
<td>2</td>
<td>SD × 24 mo</td>
</tr>
<tr>
<td>8</td>
<td>58</td>
<td>F</td>
<td>κ</td>
<td>VAD, melphalan/ASCT, XRT</td>
<td>IL-2</td>
<td>2.5 yr</td>
<td>Yes</td>
<td>3</td>
<td>92</td>
<td>1.6 × 104</td>
<td>1</td>
<td>SD × 6 wk</td>
</tr>
<tr>
<td>11</td>
<td>54</td>
<td>M</td>
<td>κ</td>
<td>VD, melphalan/ASCT, VDT-PC, VDT-AC</td>
<td>IL-7/IL-15</td>
<td>5 d</td>
<td>No</td>
<td>3</td>
<td>91</td>
<td>1.8 × 104</td>
<td>1</td>
<td>NR</td>
</tr>
<tr>
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<td>60</td>
<td>M</td>
<td>κ</td>
<td>RD, VD, VAD, VDT-PC, pomalidomide, LGH447</td>
<td>IL-7/IL-15</td>
<td>4 wk</td>
<td>No</td>
<td>3</td>
<td>91</td>
<td>1.8 × 104</td>
<td>1</td>
<td>NR</td>
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<tr>
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<td>69</td>
<td>F</td>
<td>IgGκ</td>
<td>VCD, melphalan/ASCT + thalidomide, DT, CyBorD, bendamustine, carfilzomb, carfilzomb/lenalidomide, CTX</td>
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<td>15 wk</td>
<td>Yes</td>
<td>3</td>
<td>90</td>
<td>1.9 × 104</td>
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<td>SD × 6 wk</td>
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<tr>
<td>16</td>
<td>43</td>
<td>F</td>
<td>IgGκ</td>
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<td>IL-7/IL-15</td>
<td>19 d</td>
<td>No</td>
<td>3</td>
<td>93</td>
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<td>1</td>
<td>NR</td>
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<td>F</td>
<td>κ</td>
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<td>IL-7/IL-15</td>
<td>28 wk</td>
<td>Yes</td>
<td>3</td>
<td>94</td>
<td>1.8 × 104</td>
<td>2</td>
<td>SD × 6 wk</td>
</tr>
</tbody>
</table>

*Low-dose CTX (12.5 mg/kg). This patient was re-enrolled in the trial. (R)(V)D, (lenalidomide), (bortezomib), dexamethasone; CyBorD, cyclophosphamide, bortezomib, dexamethasone; VAD, bortezomib, doxorubicin, dexamethasone; (V)DT, (bortezomib), dexamethasone, thalidomide; PACE, cisplatin, doxorubicin, dexamethasone; ESHAP, etoposide, methylprednisolone, cytarabine, cisplatin; NR, no response.
routing B cells at the time of infusion (6% ± 4.9%), we observed an overall 50% reduction in the number of B cells within the first 2 weeks after infusion (3.2% ± 2.5%) (Figure 3A), with a transient preferential elimination of κ B cells, as documented by the decrease in the κ/λ ratio (P = 0.009) (Figure 3A) and the reduced mean fluorescence intensity of κ light chain expression (Figure 3B) during the first 2 weeks after infusion.

Although no patient had symptoms consistent with severe cytokine release syndrome (CRS), because CARTs have been associated with severe CRS, we nonetheless measured cytokine levels serially in serum samples. Although we were able to detect elevation of inflammatory cytokines, such as IL-6 (Figure 3C and Supplemental Figure 3), which generally coincided with peak CART expansion, the fold increase from baseline (20-fold average) was much lower than that observed in patients who experience severe CRS (usually >75-fold) (3).

We investigated whether the treated patients developed an Ab response to the transgene. Human anti-mouse Ab (HAMA) levels almost always remained below the threshold of detection and always below levels likely to be biologically active (maximum level 12.5 U/ml; a negative HAMA assay is defined as ≤74 U/ml) (Figure 3D). Tregs were also monitored and, although we were able to detect expansion or persistence of CARTs or with clinical response, had no response to k.CARTs (Figure 5). Seven weeks later, however, he had recurrence of disease (in the axillary lymph node) despite a fourth infusion. His disease has, however, remained with minimal progression for more than 1 year.

We further analyzed the antitumor immune responses in the 3 who had received a final cycle of chemotherapy within 4 weeks of k.CART infusion, followed by a second infusion 8 weeks later, again with transient SD, the patient received 1 dose of bendamustine because of active disease. Four weeks later, he had evidence of minimal residual disease in the neck by PET/CT scan. He received one k.CART infusion, followed by a second infusion 8 weeks later, without intervening chemotherapy. The patient entered a CR after the second infusion, and this response has been sustained for almost 3 years (Figure 4). Patient 9 had progressive improvement of pelvic lymphadenopathy and achieved a CR after the third infusion of k.CARTs (Figure 5). Seven weeks later, however, he had recurrence of disease (in the axillary lymph node) despite a fourth infusion. His disease has, however, remained with minimal progression for more than 1 year. Patient 6 (lymphoplasmacytic lymphoma [LPL]) and patient 11 (CLL) had transient PR and SD, respectively. The remaining 5 of 9 NHL or CLL patients (including the 3 who had received a final cycle of chemotherapy within 4 weeks of k.CART infusion) had PD.

Four of seven MM patients had responses. Patients 4 and 7 had prolonged SD for 17 and 24 months. Patient 4 had only minimal residual disease (MRD) after high-dose melphalan and autologous stem cell transplantation (ASCT) (positive immunofixation with unmeasurable monoclonal protein) and maintained stable MRD for 17 months after a single k.CART infusion. Patient 7 had a progressive decline in her paraprotein levels (2,240 to 1,730 mg/dl IgG) and improvement in anemia (11.6 to 14.4 g/dl hemoglobin), which was sustained for 2 years (Figure 6). Neither of these 2 patients was on lenalidomide (or any other immunomodulatory drugs) at the time of k.CART infusion, either because they had previously progressed on the drug or were intolerant of it. Two other patients (nos. 8 and 14) met the criteria for transient SD, namely a less than 50% reduction and a less than 25% increase in paraprotein levels (38% free κ light chain reduction in patient 8 and 11% increase in IgGκ in patient 14). Patient 8 was re-treated with k.CARTs 1.5 years later, after receiving additional conventional therapy, again with transient SD. Three of seven patients, including the two who had received a final cycle of chemotherapy within 4 weeks of k.CART infusion, had no response to k.CARTs.

We investigated whether the patients’ responses correlated with numbers of k.CARTs infused, blood cell counts at infusion (total white blood cell, absolute lymphocyte, and absolute neutrophil counts), circulating Ig and κ light chain levels, and plasma cytokine levels. We found no significant association between these parameters and response.

We further analyzed the antitumor immune responses in the recipients to seek evidence for newly emergent immunity to additional tumor-associated antigens. No evidence for such “epitope spreading” was obtained, as we found no differences in the fre-
with diffuse large B cell lymphoma (DLBCL), the rates of CRs to CD19.CARTs correlate with the intensity of the preconditioning chemotherapy before CD19.CART infusion, ranging from 12.5% (17) to 57% (12), according to the doses of the lymphodepleting drugs fludarabine and CTX. Our patients received no or limited lymphodepleting chemotherapy (12.5 mg/kg CTX) that produced a maximum 30% reduction in lymphocyte counts: indeed, the 2 patients who had a CR received no pretreatment lymphodepletion. More profound lymphodepletion immediately before κ.CART infusion, by depleting Tregs, downregulating immunoinhibitory pathways such indoleamine 2,3-dioxygenase (IDO) (18) in the tumor environment, and creating a favorable homeostatic cytokine milieu for T cell expansion, will likely increase the therapeutic activity of κ.CARTs.

Our clinical study also enrolled patients with MM. There are few reported clinical data for MM patients treated with CARTs. A recent report from China describes the outcomes of 5 MM patients treated with CD138.CARTs, 4 of whom achieved up to 7 months of SD (19). We observed modest antimyeloma effects, documented by a reduction of the paraprotein, a selective reduction of free κ light chains, and an improvement of anemia. These effects were sustained for up to 24 months.

Discussion

We report the first clinical trial to our knowledge involving the treatment of B cell–derived malignancies with CARTs targeting a clonally restricted B cell marker intended to limit B cell aplasia while maintaining antitumor activity. We found that adoptive transfer of CARTs that target the κ light chain of Igs is feasible and safe at all dose levels studied and that the κ light chain is a validated target for NHL, since κ.CARTs can induce clinical responses, including sustained CRs, in patients with NHL.

The US Food and Drug Administration (FDA) has granted the “breakthrough” designation to CD19.CARTs for the treatment of refractory or relapsed ALL on the basis of the high rate of clinical responses achieved in different clinical studies (3, 5, 6). In contrast, the efficacy of CD19.CARTs in patients with refractory or relapsed CLL or SLL and other NHLs appears less striking. For example, the overall response rate in a study of CD19.CART treatment of CLL is currently 57% (16), while in patients with diffuse large B cell lymphoma (DLBCL), the rates of CRs to CD19.CARTs correlate with the intensity of the preconditioning chemotherapy before CD19.CART infusion, ranging from 12.5% (17) to 57% (12), according to the doses of the lymphodepleting drugs fludarabine and CTX. Our patients received no or limited lymphodepleting chemotherapy (12.5 mg/kg CTX) that produced a maximum 30% reduction in lymphocyte counts: indeed, the 2 patients who had a CR received no pretreatment lymphodepletion. More profound lymphodepletion immediately before κ.CART infusion, by depleting Tregs, downregulating immunoinhibitory pathways such indoleamine 2,3-dioxygenase (IDO) (18) in the tumor environment, and creating a favorable homeostatic cytokine milieu for T cell expansion, will likely increase the therapeutic activity of κ.CARTs.

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In preclinical studies, we did not observe deleterious effects of physiological levels of soluble Igs on the function of κ.CARTs (15). Indeed, soluble Igs did not impair the capacity of κ.CARTs to eliminate κ− tumor cells and appeared to sustain the expansion of κ.CARTs, which may partially explain the observed clinical responses to κ.CARTs in patients with NHL. However, we also cannot exclude the possibility that higher levels of free κ light chains, such as those seen in κ-restricted MM, cause adverse effects on κ.CARTs, either by blocking the recognition of κ light chains on the surface of targeted cancer cells or by causing excessive T cell stimulation and exhaustion, thus limiting the therapeutic effect. On the other hand, plasma cells, including malignant counterparts, have minimal (if any) surface expression of light chains, and that can explain, to some extent, why the clinical responses were less striking in patients with MM. It would have been interesting to measure the distribution and persistence of infused κ.CARTs in the bone marrow of patients with MM, but the IRB-approved protocol did not mandate bone marrow studies as long as there was measurable disease by paraproteinemia, which was present in all of these patients.

How can we improve upon these outcomes? Clinical data in patients with ALL infused with CD19.CARTs suggest that the nature of the costimulation provided within the CD19.CAR may affect CART in vivo persistence. Indeed, while CD28 and 4-1BB costimulation of CD19.CARTs are equally effective in inducing clinical responses in patients with relapsed ALL, 4-1BB costimulation may ensure a longer persistence of CARTs and potentially more durable remissions (5). In addition, early in vitro studies and more recent animal experiments suggest that CARs that include long spacers derived from the constant regions of human Igs may...
We conducted a phase 1 study (ClinicalTrials.gov [36x308]. Methods [36x440] to assess the feasibility and safety of infused escalating doses of autologous, polyclonally activated peripheral blood T cells that were genetically modified to express a light chain–specific CAR (κ.CAR) in patients with relapsed or refractory κ-restricted lymphoproliferative disorders (NHL, CLL, and MM). All patients had to have measurable disease at the time of infusion (Figure 1). Patients with CLL or an ALC above 500/μl and who had not received any chemotherapy within the previous 4 weeks were given low-dose (12.5 mg/kg) CTX as a single infusion 4 days before administration of the first dose of κ.CARTs. Patients on chemotherapy at the time of treatment could receive κ.CARTs no sooner than 4 days after finishing their last cycle of chemotherapy. Otherwise, no chemotherapy was given to patients before or after κ.CART infusion. We administered 3 dose levels of κ.CARTs: 2 × 10^7 (dose level 1); 1 × 10^8 (dose level 2); and 2 × 10^8 (dose level 3) cells/m². These doses were based on total cell numbers, regardless of the percentage of CAR⁺ cells. We used an interpatient dose escalation protocol that followed a continual reassessment method, which required safety to be demonstrated 6 weeks after infusion in 2 patients at each dose level before escalation. Additional infusions of the same dose of κ.CARTs were allowed as long as there was evidence of clinical benefit, defined as at least SD.

Additional infusions were administered at least 6 weeks apart, and no chemotherapy was given before κ.CART infusion. Clinical and laboratory evaluations were performed at weeks 1, 2, 4, and 6 and months 3, 6, 9, and 12 after CART infusion, and yearly thereafter. Adverse events during and after T cell infusions were graded according to the NIH Common Terminology Criteria for Adverse Events (CTCAE), version 4 (http://ctep.cancer.gov). Responses were assessed by imaging or laboratory studies, as applicable, at week 6 after CART infusion and were defined as CR, PR, SD, or PD, according to the NCI’s Response Evaluation Criteria in Solid Tumors (RECIST) (23), with modifications based on the International Working Group criteria for NHL (24); the National Cancer Institute Working Group for CLL (25); or the modified (International Myeloma Working Group [IMWG]) uniform response criteria for MM (26).

Generation of retroviral constructs. We cloned the Ab targeting the κ light chain of human Ig produced by the CRL-1758 mouse hybridoma (ATCC) as a single chain (scFv) and generated a second-generation CAR construct containing both CD28 and ζ endodomains (15). A spacer region derived from the human IgG1-CH2CH3 domains was cloned in-frame between the scFv and the signaling domains to facilitate detection by phenotypic analysis of the transgenic product. This cassette was then cloned into the SFG retroviral backbone. We generated the clinical-grade packaging cell line using PG13 cells (gibbon ape leukemia virus pseudotyping packaging cell line, CRL-10686; ATCC) (27). We used the highest-titer clone to establish a master cell bank and released the clone for clinical use only after safety testing and vector sequencing. We stored the final viral supernatant at −80°C and tested it before clinical release.

Generation and transduction of κ.CARTs. To generate κ.CARTs, we transduced PBMCs obtained by Ficoll density centrifugation of peripheral blood obtained by phlebotomy 1–3 months before CART administration. We activated PBMCs with OKT3 Ab (Janssen Biotech) or CD3 and CD28 Abs (Miltenyi Biotec) and recombinant human IL-2 (IL-2, 100 U/ml; Prometheus) or IL-7 (10 ng/ml; R&D Systems) and IL-15 (5 ng/ml; R&D Systems) in 24-well plates precoated with a recombinant fibronectin fragment (FN CH-296, Retronectin Takara; Clontech). After transduction, T cells were expanded ex vivo in the presence of recombinant human IL-2 (rhIL-2) (100 U/ml) or IL-7 (10 ng/ml) and IL-15 (5 ng/ml), added twice a week. A change to the process of manufacture took place during the trial, when IL-7 and IL-15 replaced IL-2 because of our own and others’ observation that the combination of IL-7 and IL-15 better preserves the CD8⁺CD45RA⁺CCR7⁺ subset in culture, which is associated with superior expansion and persistence of CARTs in vivo (28). The characteristics of the T cell products are summarized in Supplemental Figure 1 and Supplemental Table 1. The generation of κ.CARTs required a median of 17 days (range 10–27) of culture in the 14 T cell products generated using IL-2 and 16 days (range 12–17 days) in the 13 T cell products expanded in the presence of IL-7 and IL-15. The expansion of T cells grown in IL-7 and IL-15 was overall better than that of T cells grown in IL-2 (Supplemental Figure 1A).

Immunophenotyping. We used phycoerythrin (PE), FITC, peridinin chlorophyll protein (PerCP) or allophycocyanin-conjugated (APC-conjugated) CD3 (clone SK7); CD4 (SK3); CD8 (SK1); CD56 (B159); CD19 (J25Cl); TCR-αβ (T10B9); TCR-γδ (B1); CD62L (DREG56); CD27 (M-T271); CD28 (CD28.2); CD45RA (H1100); CD45RO (UCHL1); CCR4 (48607); CCR4 (1G1); CCR5 (2D7/CCR5); CCR7 (3D12); CXCR3 (1G6/CXCR3); CXCR4 (1G5); CD162 (KPL-1); CD54 (HA58);
CD38 (H7B); CD106 (51-10C9); CD11a (HI11); CD11c (B-ly6); CD18 (6.7); and κ (TB28-2) and λ (I-155-2) light chain Abs (all from BD Pharmingen) to stain the T cell products or tumor cells. We included control samples labeled with the appropriate isotype-matched Abs in each experiment. The κ.CAR was detected with an Fc-specific Alexa Fluor 647-conjugated goat polyclonal F(ab′)2 fragment anti-human IgG (catalog 109-606-088; Jackson ImmunoResearch), which recognizes the IgG1-CH2CH3 component of the CAR. Cells were analyzed with a FACScan (BD) equipped with a filter set for 4 fluorescence signals, using CellQuest software. Data analysis was performed using FlowJo Software (Tree Star). In selected experiments, frozen PBMCs from pre-and post-CART infusion time points were thawed and the number of CD19+ cells and of κ+ cells enumerated by FACS using CountBright beads (Invitrogen).

Cytotoxicity assays. The cytotoxic specificity of each T cell line was measured in a standard 4-hour ⁵¹Cr release assay using effector-to-target (E:T) ratios of 40:1, 20:1, 10:1, and 5:1. Daudi (κ+ tumor cells), HDLM-2 (κ+ tumor cells), and K562 (NK-sensitive cells) were used as target cells. The target cells were labeled simultaneously for 1 hour with ⁵¹Cr. We calculated the percentage of specific lysis as follows: [(experimental release – spontaneous release)/(maximum release – spontaneous release)] × 100.

Real-time quantitative PCR of the κ.CAR transgene. We quantified the integrated genome of the retrovirus encoding the κ.CAR by real-time quantitative PCR (qPCR) (13). After extracting DNA from peripheral blood samples with the QIAamp DNA Blood Mini Kit (QIAGEN) following the manufacturer’s instructions, the DNA was amplified with primers and probes (Applied Biosystems) complementary to specific sequences within the retroviral vector (15). The standard curve was established using serial dilutions of the plasmid encoding the transgene. Amplifications were performed using the ABI 7900HT Fast Real-Time PCR System (Applied Biosystems, Thermo Fisher Scientific) according to the manufacturer’s instructions.

Cytometric bead array and multiplex assays. We analyzed plasma or serum samples collected before and after κ.CART infusion using a BD cytokine cytometric bead array (CBA) kit (BD Pharmingen) or a Milliplex Kit (EMD Millipore), following the manufacturers’ instructions. In parallel with the samples, we used the human cytokine standards provided with the kits to prepare standard curves. The CBA assays were run using the BD FACSCalibur, and the data were analyzed from the Milliplex kits using the Luminex 200 System and Milliplex Analyst Software (EMD Millipore).

ELISPot assays. IFN-γ enzyme-linked immunospot (ELISpot) assays were performed to identify precursor T cells specific for known TAAs. PBMCs were plated at 2 × 10⁸ to 4 × 10⁹ per well in triplicate in PI-Multiscreen plates (EMD Millipore), and IFN-γ production was measured in response to stimulation with peptide mixtures (pepmixes) consisting of 15-mer peptides overlapping by 11 aa and spanning the entire protein of interest for the following antigens: pp65, NY-ESO-1, MAGEA4, PRAME, and P53 (0.6 nmol of each peptide) (JPT Technolog-}

6. Lee DW, et al. T cells expressing CD19 chi-


