Method articles

Chimeric antigen receptor preparation from hybridoma to T-cell expression

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Abstract

The successful use of chimeric antigen receptor (CAR) for hematological cancer treatment has influenced the direction taken in translational research toward an increasing focus on personalized targeted immunotherapy. Thus, a growing number of labs worldwide are now interested in testing their old antibody collections in this format to broaden the spectrum of utility and improve safety and efficacy. We herein present a straightforward protocol for the identification of an antibody from a hybridoma and the design of the single chain fragment that will be placed on the extracellular part of the CAR construct. We further show how to test the expression and the activity of the construct in primary T cells. We illustrate our demonstration with two new CARs targeted against the B cell receptor, more precisely the light chains κ and λ, that represent potential alternatives to the CD19 CAR used in the treatment of B-cell malignancies.

Statement of Significance: Chimeric antigen receptor molecules are becoming increasingly popular, and an easy-to-follow method to isolate and express them is herein presented.

KEYWORDS: hybridoma; antibody; single chain variable fragment; chimeric antigen receptor

INTRODUCTION

Chimeric antigen receptor (CAR)-expressing T cells are now used to treat leukemia and lymphoma patients who are refractory to standard therapy. These powerful cells combine the specific target recognition offered by an antibody fused with parts of the natural signaling machinery of the T-cell receptor (TCR) (Fig. 1A). CAR is usually expressed in immune effector cells (NK or T cells) isolated from the patients (autologous settings), from a donor (allogeneic settings) or in a cell line. They are genetically transformed ex vivo or in vitro, (re)-injected into the patients and expected to reach the tumor and kill it [1].

A CAR molecule at the surface of an effector cell will, upon target recognition, form clusters and trigger endogenous signaling component recruitment. The intracellular signaling tail is a combination of domains that generate a fast response in the effector cell against the target (killing and cytokine release) and a long persistence signal to allow a durable and lasting response. Thus, a CAR is able to transform a polyclonal population of effector cells into tumor killing cells by retargeting them against a selected target. This is what was successfully achieved when patients were treated with anti-CD19 CAR T cells [2]. Here, a heterogeneous population of T cells was redirected to gain specificity against the B-cell marker CD19, which lead to the destruction of malignant B cells as well as the healthy ones. The latter has been shown to lead to complications in patients and stimulated the quest for alternative antigens, such as the ones we present here, targeting only a subgroup of the population.

Although antibodies have been sequenced for almost four decades [3] and a very comprehensive protocol was released 20 years ago to isolate antibody sequences [4], a practical protocol for the design of CAR molecules does...
not seem to be available. We herein present a method that can be adapted in any lab to achieve the creation of a CAR. We illustrate it by describing the isolation of the coding sequence from two validated hybridomas producing antibodies with specificity against the B cell receptor (BCR) light chain -Igκ and -Igλ chains. We will explain how the single chain variable fragment (scFv) sequence was extracted and designed to create the binding part of the CAR. Finally, we will also give insight into the preliminary tests that should be run to confirm the expression and activity of a new CAR. The complete procedure is displayed in Fig. 1B.

**MATERIALS AND METHODS**

**5'-RACE**

*mRNA preparation.* Pellets from 1 million cells and isolation of the total RNA were done with a starting volume of 600 μL lysis buffer, then every step of the manufacturer’s protocol was followed (Absolutely RNA Miniprep, Agilent Technologies, USA) including the DNAse I treatment. Importantly, the lysis of each pellet was divided in two and the protocol was performed in parallel to ensure that somatic mutations detected are original and not coming from the use of a Taq polymerase. Final elution
was performed in 50 μL of pre-warmed buffer. All mRNAs were quality/quantity checked on a Nanodrop apparatus (Thermo Fisher, USA) and stored at −20°C until cDNA preparation.

cDNA synthesis. The reverse transcription was performed using SuperScript III Reverse Transcriptase (200 U/μL, Invitrogen, Eureka, CA, USA) with a reaction mix of $V_{\text{final}} = 20.5 \text{ μL}$, prepared in two parts containing: 1 μL oligo-dT (50 μM, Invitrogen), 1 μL dNTPs (10 mM, Thermo Fisher), 1 μg mRNA and dH2O to a final V of 14 μL. This part was then incubated at 65°C for 5 minutes and chilled on ice, then the rest of the mix was added as follows: 4 μL first strand buffer, 1 μL DTT (0.1 M, Invitrogen), 0.5 μL RNasin (40 U/μL, Promega) and 1 μL Reverse Transcriptase. The mix was incubated at 50°C for 1 hour, then the RNasin was inactivated by 15 minutes incubation at 60°C. The mix was treated with 1 μL RNaseH (5000 U/mL, New England Biolabs, USA) and 1 μL TdT incubated for 15 minutes at 37°C for 20 minutes, to remove the remaining mRNA. The nucleotides were precipitated by NaAc-EtOH method: 21.5 μL cDNA is mixed with 0.5 μL Glycogen (20 mg/ml, Thermo Fisher), which is used as carrier, and 1/10 V of NaAc (3 M pH 5.5, Invitrogen) and 2.5 V EtOH 100%, incubated at −20°C for minimum 30 minutes. The precipitated cDNA was spun down at 10,000g at 4°C for 10 minutes, washed once with 200 μL 70% EtOH and finally resuspended in 21 μL dH2O. One μL was used to control the presence of product on an agarose gel, 10 μL for dC tailing and 10 μL kept at −20°C.

5′dC-tailing. Samples were placed at 95°C for 1 minute and directly chilled on ice; this will break the structures formed in the DNA that might mask the site of tailing. The reaction was performed using TdT enzyme and buffer mix (400 U/μL, Roche, Switzerland): 10 μL cDNA, 4 μL TdT reaction buffer 5X, 4 μL CoCl2, 1 μL dCTP (10 mM, Thermo Fisher) and 1 μL TdT incubated for 15 minutes at 37°C, and the reaction was stopped by adding the NaAc-EtOH precipitation components as described in the previous section.

Nested PCR. One microliter of the dC-tailed cDNA was finally amplified in a serial nested PCR using the constant part-specific primers and a universal dC-annealing primer (Fig. 2B). The primers were designed in-house except pGI15_TOPO [5], mlgG2bCH1, mlgGKCreV, mlgG1CH1−2, mlgG2bCH1−2 [6], mlgG2aCH1−2 [7] and mlgGKCreV-2 [8]. Each PCR was run in these conditions: 4′ 94°C for the initial denaturation and 25 cycles (1′ 94°C, 1′ 53°C, 1′ 72°C) with the following mix: 5 μL 10X Taq Buffer, 1.5 μL of primer (10 mM stock), 1 μL dC-tailed cDNA, 1 μL Taq DNA polymerase (Hospital collection) and dH2O added to a $V_{\text{final}} = 50 \text{ μL}$. Both PCR products were separated on a gel; the second is the same as the first, but 1 μL of the first reaction was used as a template. The expected size should be larger than 450 bp, a smear should be visible (Fig. 2C).

Sequence identification, scFv design and subcloning into a signaling cassette

The PCR products were run on gel, and bands were extracted and purified using NucleoSpin® Gel and PCR Clean-up kit (Macherey-Nagel, Germany); after checking the presence of the band and estimating its concentration on a gel, it was subcloned into pGEM T/A cloning vector following manufacturer’s protocol (Promega, USA) and transformed into NEB 5-alpha competent cells (New England Biolabs, USA) and blue/white colonies were screened. Single colonies were picked, and the presence of an insert was determined using EcoRI (New England Biolabs). Insert-containing vectors were sent to sequencing (Eurofins MWG, Germany) using T7 and M13(−29) primers. The sequences were analyzed using the IMGT (http://www.imgt.org/IMGT_vquest/vquest) online tool to identify the V chains, hypervariable domains and the somatic mutations. The beginning and the end of the V domain were defined according to IMGT nomenclature, and these limits were used to design the scFv. The resulting protein sequence of the construct was constructed as follows, signal sequence (METDTLLLWVLLLWVPGSTG)-VL-(G3S)4(GGGGSGGGGSGGGGGS)-VH; it was ordered as codon optimized for production in human cells (Eurofins MWG), and restriction sites were added (NcoI/BamHI) to subclone it into a signaling tail-containing vector (pENTR-seq8843). Verified clones were recombined using the Gateway cloning system (Invitrogen) into pMP71 converted plasmid [5].

T-cell transduction, CAR detection and killing assay

T-cell transduction, cytokine staining and bioluminescence (BLI)-based killing assay have been described in details elsewhere [9]. Briefly, primary T-cell transduction with retroviral particles was performed by double spinoculation followed by expansion on CD3/CD28 beads. The staining of the CAR was performed with Biotin-SP-AffiniPure F(ab)2 Fragment Goat Anti-Mouse IgG (Jackson ImmunoResearch, USA), followed by a secondary staining with Streptavidin-PE (Biolegend, USA). CAR-expressing T cells were co-cultured with cells positive or negative for the target antigen, and TNF-α was detected by flow cytometry using TNF-α-PE (MAb11, BD Biosciences). The samples were run on a BD FACSauto flow cytometer (BD Biosciences, USA), and data were analyzed using FlowJo software (Treestar Inc., USA). BLI killing assay was also based on a co-culture experiment where target cells permanently expressed firefly Luciferase gene, and killing was correlated to the luminescence signal [10].

Expression validation of the target cells

Target cells were assessed for their expression status for Igκ and Igλ by flow cytometry. The first assessment was done by staining with the following commercial antibodies; anti-human Igκ-APC (MHK-49, Biolegend) and anti-human Igλ-PE (1-155-2, Thermo Fisher). The same cell lines were also stained with the hybridoma supernatants with varying dilutions, followed by a secondary staining with anti-mouse IgG-APC (Poly4053, Biolegend).
Figure 2. Investigating the hybridoma by 5′-RACE. (A) The depiction of 5′-RACE step by step protocol. (B) List of primers used in this study and references to already published ones. (C) Acquired gel pictures throughout the 5′-RACE protocol, each kappa and lambda hybridomas were divided into two and the protocol was followed in two replicates (e.g., named as K#1 and K#2). The top picture shows the result of the cDNA synthesis, the middle pictures the first PCR results of dC tailed samples and the bottom pictures the result of the second PCR performed with the first PCR products. The ladder information corresponding to 500 and 400 bp bands were specified as well as the primer dimer with an asterisk on each gel image.
Statistical analysis

Student’s t-test was used when two groups were compared. BLI killing assay was analyzed by non-linear regression (curve fit) on GraphPad Prism® (GraphPad Software, USA).

RESULTS

Isolation of the antibody coding sequence from a hybridoma

We previously created hybridomas producing anti-human Igκ and Igλ antibodies (clones K13 and K16, respectively). The cells were grown and supernatants were tested for antibody production and confirmation of the target specificity, as shown in Fig. 1C; we compared the target recognition of our antibodies to commercial ones. We distinguished the BCR light chain isoforms by using model cell lines known to be positive for either κ or λ.

When the hybridoma culture reached around 1 million cells, pellets were prepared, lysed in mRNA extraction buffer and frozen until use (see Materials and Methods for kit listing and detailed procedure). Fig. 2A depicts the whole cloning procedure: first total mRNA was used to synthesize cDNA using oligo-dT as a primer. After treatment with RNaseH to digest remaining mRNA, the cDNA was dC tailed on its 5’-end using TdT enzyme. The reaction was stopped after 15 minutes and the tailed cDNA extracted by precipitation. This product was checked on a gel (Fig. 2C, top panel) and used as a template for a nested PCR using G1 oligo that binds the TdT synthesized C-tail and antibody specific primers (Fig. 2B) annealing to the constant domain. The first and second amplifications were run on a gel (Fig. 2C, middle and bottom panels, respectively). These PCR reactions gave a first indication on the identity of the antibody chain type. As shown both anti-Igλ and anti-Igκ possess IgG1 as heavy chain and Igκ as light chain. Since IgG1 and IgG2a sequences are related we can observe their amplification with the different sets of primers; however the sequence obtained after sequencing of the antibody will provide information on the exact IgG identity.

The PCR reactions were then subcloned: since the PCR reactions were conducted with a Taq polymerase, one-nucleotide overhang was created and could accommodate a T/A cloning system. Bacterial colonies were picked and minipreps digested to estimate the size of the insert that is expected to be higher than 450 bp, but might vary due to the tailing of differently sized mRNA. This range ensures that a piece of the 5′-UTR will be present; hence the natural start detected (Fig. 3A).

Identification of V-chain types and scFv design

The sequences were then identified using the IMGT database. In addition, somatic hypermutations were identified and the limit of the V chains was defined, which facilitates the further design of the V chains in the scFv (Fig. 3A). The preparation of scFv can be designed in multiple manners, but we experienced that the design starting with the light chain variable fragment (VL) including a secretion sequence (L chain) linked to the variable heavy chain (VL) by glycine serine repeats (G₄S₄)₄ was normally adaptable to all our constructs. Nevertheless, it is always an advantage to compare different designs by swapping VL and VH or testing linkers of different size if the “classical” design is not working or with low efficiency [11].

The protein sequence of the scFv fragments of the two antibodies was then ordered as a synthetic DNA, which was subsequently cloned in our system. Here the second-generation signaling tail was used (Fig. 2B). The final product of each CAR construct will thus be L-VL-linker-VH-CD8hinge-CD8TM-4-1BB-CD3ζ. The sequences of the different gene blocks are depicted under each entity and were based upon description in Uniprot database: CD8hinge and CD8TM, CD8A_HUMAN (P01732), 4-1BB, TNR9_HUMAN (Q07011), CD3ζ and CD3Z_HUMAN (P20963).

CAR expression and activity detection

The complete CAR coding sequence was then subcloned into a retroviral expression vector and particles were prepared using Gateway-compatible system as we previously described [5]. We routinely test the expression of our constructs in an easily manipulated cell line such as Jurkat cells before proceeding to primary cells. Indeed, this cell line can be transduced in one spinoculation and the transferred gene
can usually be detected within 2 to 3 days. Fig. 4A shows the expression of the CARs in Jurkat cells detected by flow cytometry using specific antibodies against mouse antigen-binding fragment (Fab). From these data we confirmed that the design was leading to a correctly folded product expressed at the cell surface. We further validated this result by using a system based on primary human T cells. Here, stimulated T cells isolated from buffy coat were used as effector cells and transduced by double spinoculation (see Materials and Methods). Seven days after spinoculation, T cells were tested for CAR expression (Fig. 4B).

Analysis of CAR reactivity was performed using primary T cells in co-culture assay with target cells expressing the cognate antigen. Here BL-41 and MAVER-1 cell lines were used since their BCRs are Igκ+ and Igλ+, respectively. The target cells were incubated with CAR T cells or mock-transduced T cells. The latter are used to set the baseline of the stimulation in order to distinguish specific CAR response from background. Indeed, T cells from diverse donors might react differentially against various targets; this can be explained by a difference in HLA composition between the blood donor and the target cells.
DISCUSSION

In this study we introduce a complete protocol for CAR preparation. We depicted the details presently being tested to conveniently extract an antibody sequence from a hybridoma and further design from this sequence a molecule that will redirect effector immune cells. We then displayed the minimal experiments required to validate that a given construct turns effector cells into a tumor cell killer. It is worth saying that due to the growing interest for using human antibodies as a source for CAR constructs, this protocol could be adapted. If humanized mice are used [14], only the primer sequences should be changed to human specific ones.

The clinical success of CD19 CAR in the treatment of B-cell malignancies has demonstrated the power of immunotherapy [2]. However, recent clinical data have shown that alternative solutions to CD19 CAR are needed; some patients become refractory to the treatment through several mechanisms directly related to the target recognition or the CAR itself [15]. Alternative B-cell CAR constructs have been identified [16–18] and some are presently being tested in the clinic [19, 20], even in alternative formats such as scFv placed in tandem on a single construct [19]. Importantly, there is growing evidence that more than one CAR molecule should be tested to identify the CAR with optimal affinity and function. Even if the original antibody has a high affinity for the target and the in vitro testing of the CAR construct is successful, the in vivo assessment might not follow this trend [21], thus more hybridomas will need to be tested. Although some of these CARs such as the CD22 CAR clone m971 [16] were isolated from phage display libraries [22], the majority of molecules still come from hybridoma, and we believe that the protocol proposed herein will help scientists to deliver the future clinical CARs.

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