Review

Engineering autonomous closed-loop designer cells for disease therapy

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SUMMARY

Synthetic biology has made it possible to engineer mammalian cells for on-demand delivery of therapeutic agents, providing therapeutic solutions for chronic or intractable diseases. Currently, most of the engineered therapeutic cells are regulated by the administration of exogenous inducers, but the need for repeated administration of these xenobiotics is problematic from the viewpoints of patients’ compliance and quality of life, as well as possible side effects. Recently, synthetic biologists started to address these issues by constructing autonomous, closed-loop therapeutic cells, often referred to as designer cells. These cells are equipped with sensing modules that detect and link marker(s) of the target disease to signaling cascades that stimulate the secretion of specified therapeutic agents in a timely and quantitative manner, without the need of exogenous inducers. Here, we review recent work on designer cell engineering and their in vivo therapeutic applications, focusing on the molecular mechanisms and signaling pathways employed.

INTRODUCTION

Cell-based therapy is opening up the possibility of novel therapeutic solutions for diseases such as organ failure and tissue dysfunction, where conventional drugs are of limited efficacy or impose a severe burden on the patients’ quality of life. For example, cell-based alternatives to repetitive and invasive drug injections in patients with chronic diseases, such as diabetes, have been extensively investigated. Indeed, in-vivo implantation/injection of healthy or genetically modified cells has been considered to treat a range of diseases, including metabolic disorders, neurodegenerative diseases, genetic defects, and multiple types of malignancies (Buzhor et al., 2014; Filipovich, 2008; Kim and Keating, 2019; Yasuhara et al., 2017; Grupp et al., 2013). To date, the best-known clinical application is probably the use of chimeric antigen receptor (CAR)-T-cells to treat various types of blood cancers that are frequently refractory to standard chemotherapy (Grupp et al., 2013; Till et al., 2008).

Early work was focused on stem cell biology, based on the discovery of pluripotent stem cells (PSCs) by James Thomson and the later introduction of induced pluripotent stem cell (iPSC) technology by Shinya Yamanaka and John B. Gurdon (Thomson et al., 1998; Takahashi and Yamanaka, 2006). However, the clinical applicability of therapies based on stem cells has been severely limited by the risk of tumor formation because of undifferentiated or incompletely differentiated stem cells (Ota, 2008; Bailey, 2012; Kimbel and Lanza, 2015; Einhorn and Iorio, 2017). For this reason, alternative strategies based on mammalian cells equipped with synthetic or modified genetic circuits and signaling networks have recently received more attention. In this approach, somatic cells are ex vivo engineered with a set of genes which enable biochemical linking of certain signal stimuli to a desired cellular output, such as the initiation of therapeutic protein secretion (Kojima et al., 2020; Scheller and Fussenegger, 2019; Heng et al., 2015) (Figure 1A). The engineered cells are then encapsulated with a semipermeable polymer to protect them from the host immune response, while allowing transfer of nutrients and other molecules (Figure 1B). Several types of immune-compatible polymers have recently been synthesized to provide in vivo-implanted cells with a protective and supportive shield suitable for long-term application (Rokstad et al., 2002; Ashimova et al., 2019). Following ex vivo encapsulation, the capsule device is introduced into the body, where the synthetically programmed cells mediate a specific therapeutic or diagnostic function (Figure 1C) (Sedlmayer et al., 2018; Teixeira and Fussenegger, 2017, 2019; Schukur and Fussenegger, 2016).

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The earliest therapeutic cells were based on controllable, open-loop gene circuits, which require exogenous signaling initiators to switch the cells’ behavior (Figure 2A). Various chemicals, heat, physical force, light, electricity, and even radio waves have been used as inducers (Figure 2B) (Doshi et al., 2020; Stefanov et al., 2021; Strittmatter et al., 2021; Müller et al., 2015; Krawczyk et al., 2020; Stanley et al., 2012). Although engineering open-loop systems is relatively straightforward, these systems have various drawbacks in a therapeutic setting. First, they rely on frequent delivery of the external inducer. Second, the functional behavior of open-loop systems is not readily predictable in vivo because of interindividual variability of physiological and pathological factors. Thus, tedious personal dosage adjustment for each patient may be required (Delco et al., 2005; Verbeeck and Musuamba, 2009). In addition, light-controlled gene circuits can be problematic because of the limited penetration of light through living tissues (Ye and Fussenegger, 2019). However, the functional dependency of the open-loop expression systems on inducer availability in the cellular environment can be advantageous in certain circumstances. For example, open-loop operating devices can be easily shut down in the event of therapeutic-induced toxicity simply by stopping the administration of the inducer. In addition, personal dosage adjustment of the activating inducer may offer an additional control point between the typical ON/OFF endpoints, providing more flexibility to achieve an optimal individual response.

Recently, there has been increasing interest in the development of genetically engineered therapeutic cells that functionally mimic specialized cells in the body (Haellman and Fussenegger, 2016). These so-called designer cells are rationally engineered to continuously and accurately sense certain physiological inputs, integrate and process this biochemical information, and respond in a programmed manner (Figure 2B). These closed-loop feedback-controlled signaling systems enable the cells to respond specifically and quantitatively to changes in disease biomarker(s) in the blood, allowing precise and timely output control. Here, we review recent work on the engineering of designer cells, as well as their therapeutic applications in in vivo disease models.
DESIGNER CELLS FOR THE TREATMENT OF METABOLIC AND ENDOCRINE DISEASES

Diabetes mellitus (DM) affects more than 45 million people worldwide (Kharroubi, 2015). Patients diagnosed with DM experience chronically high blood glucose levels, which may lead to life-threatening complications in the absence of medical intervention (Melendez-Ramirez et al., 2010; Papatheodorou et al., 2018). Type-1 DM is a consequence of autoimmune destruction of insulin-secreting pancreatic β-cells, whereas type-2 DM involves insulin resistance of the peripheral tissues (Banday et al., 2020). Therapeutically, insulin injection is the main treatment for type-1 DM and for complicated cases of type-2 DM (Silver et al., 2018). However, repeated injections are invasive and tend to reduce the patients’ compliance and quality of life. Therefore, efforts have been made to develop synthetic β-cell-mimicking designer cells. Xie et al. linked high glucose levels to insulin secretion using non-endocrine designer cells, and showed that their strategy worked in a DM mouse model (Xie et al., 2016). They utilized ATP overproduction in response to high intracellular glucose levels as the input signal (DM biomarker) (Figure 3A). Elevated levels of ATP in the cytoplasm lead to closure of the ATP-sensitive potassium channel (K\text{ATP}), inducing a transient cellular depolarization that is further amplified by the opening of voltage-dependent calcium and sodium channels, resulting in a significant increase of intracellular calcium concentration. The resulting increase of stored cytoplasmic calcium was linked to PNFAT-driven insulin transcription through nuclear translocation of a calcium-responsive transcription factor. These cells proved effective to restore glucose homeostasis in a type-1 DM mouse model. Ye et al. engineered therapeutic designer cells that sense high insulin levels and respond by producing adiponectin, which is a protein hormone that reverses insulin resistance and restores glucose homeostasis and a normal lipid profile (Ye et al., 2017). In their work, HEK-293T cells were ectopically engineered with insulin receptor (IR) and a chimeric transcription factor consisting of tetracycline repressor, TetR, fused to the human ELK1-derived transactivation domain (TetR-Elk1) (Figure 3B). When insulin binds to IR, it initiates a series of signal transduction cascades that result in the phosphorylation of the TetR-Elk1 hybrid. The phosphorylated TetR-Elk1 is then shuttled to the nucleus, where it induces adiponectin expression through a P-TetR-driven promoter. These designer cells reverse insulin resistance and prevent lipid-caused complications in an ob/ob mouse model.
Figure 3. Designer cells to treat metabolic diseases

(A) The engineering of pancreatic β-cell-mimicking designer cells. High levels of glucose cause overproduction of ATP, inducing closure of ATP-sensitive potassium channels. Accumulation of potassium ions in the cytoplasm causes mild membrane depolarization, which activates voltage-dependent calcium and sodium channels. These ion shifts lead to a high level of calcium in the cytosol, and this induces dephosphorylation of NFAT. Dephosphorylated NFAT is translocated to the nucleus to initiate gene expression. In this work, HEK-293T cells were ectopically engineered with a Cav1.3 channel and PNFAT-insulin expression cassette in order to link high glucose levels to insulin secretion.

(B) The engineering of designer cells to overcome insulin resistance in peripheral tissues. HEK-293T cells were engineered with plasmids encoding to insulin receptor (IR), TetR-EIk1 fusion protein, and adiponectin under a TetO7 promoter. High insulin levels in the blood result in insulin binding to the IR receptor on the designer cell membrane and this activates the MAPK pathway, leading to phosphorylation of the TetR-EIk1 chimeric transcription factor. The phosphorylated transcription factor is translocated to the nuclease, and induces adiponectin gene expression.

(C) The engineering of synthetic β-cell-like therapeutic machines for diabetes treatment. Liposome-based structures, called outer large vesicles (OLVs), have been chemically fabricated with GLUT2, gramicidin A, and peptide E anchored in the bilayer part. The luminal space of OLV contains soluble glucose oxidase, catalase, and insulin-loaded inner small liposomes (ISVs). Under normal conditions, the fusion of ISVs is prevented by steric hindrance involving the PEG–CDNA and GDNA–CH bulky groups. When the glucose concentration increases and glucose enters OLV through GLUT2, it is enzymatically converted to gluconic acid by glucose oxidase. The resulting high proton levels decouple PEG–CDNA GDNA–CH aggregation, which permits the formation of coiled-coil interactions between peptide K and peptide E. These strong protein-protein interactions eventually enforce IVS fusion and thereby release insulin.

(D) The development of designer cells to manage ketoacidosis. HEK-293T cells were engineered with TDAG8, a pH-sensitive G protein-coupled receptor (GPCR). High levels of hydrogen ions stimulate TDAG8 and activate the adenylate cyclase pathway. The production of the second messenger cAMP activates protein kinase A (PKA), which phosphorylates cAMP response element-binding protein (CREB). Phosphorylated CREB binds to its response element, CRE, promoting insulin transcription.

(E) Free fatty acids (FFA)-regulated designer cells for management of diet-induced obesity. The synthetic transcription factor, TtgR-PPARγ, binds to the TtgR response sequence and represses gene expression. At high concentrations, FFAs bind to TtgR-PPARγ and elicit conformational change, leading to the recruitment of cellular coactivators that facilitate pramlintide gene expression.
Totally synthetic “β-cells”, also called artificial beta cells (AbCs), have been developed as “sense-reactive” designer cells for diabetes management (Chen et al., 2018). In this work, Chen et al. chemically constructed liposome-based vesicles-in-vesicle superstructures, which are equipped with the molecular machinery needed for mimicking β-cells, by synthesizing insulin-containing inner small liposomes (ISVs), similar to insulin storage granules in mature β-cells (Figure 3C). These ISVs are encapsulated with a bigger bilayer, called the outer large vesicle (OLV), to form a synthetic plasma membrane. Glucose transporter 2 (GLUT2) is anchored to OLVs in order to allow glucose shuttling. In addition, they placed glucose oxidase and catalase in the “cytoplasmic” part of OLVs to enable the conversion of glucose molecules to gluconic acid. Overall, these reactions lead to an increase of acidity inside the intra-OLV compartment as a function of glucose level. Gramicidin A was also inserted in the OLV membrane, in order to buffer the proton concentration over time. Mechanistically, high glucose levels inside OLVs causes a dramatic decrease of pH, which induces dehybridization of the polyethylene glycol-conjugated cytosine-rich DNA (PEG–CDNA) and the cholesterol-ended guanine-rich DNA (GDNA–CH) anchored on the ISVs (shown in black and orange). The proton-induced dehybridization serves to deshield peptide K, sterically exposing it to peptide E, an anchor protein constitutively localized in the inner surface of the OLV. Coiled-coil interactions between these peptides force the fusion of ISVs and thereby release insulin. This system has been successfully tested in an in vivo mouse model, with promising results. However, despite its ability to autonomously regulate glucose levels, this device has to be administered frequently as it lacks the ability to replenish empty ISVs with new insulin cargo.

Ketoacidosis (DKA) is a serious hyperglycemic emergency in people with DM, especially type-1 DM (Dhatariya et al., 2020). This life-threatening metabolic condition is because of a drastic reduction or total loss of circulating insulin, leading to the overproduction of acidic ketone bodies and a marked decrease of blood pH (Rosival, 2013). The most effective treatment of this rapidly deteriorating metabolic disturbance is insulin injection. Ausländer et al. developed designer HEK-293T cells, called HEK<sub>pH-Guard</sub>, which could effectively reverse KDA in a type-1 DM mouse model (Ausländer et al., 2014). HEK<sub>pH-Guard</sub> Cells are engineered to express TDAG8, which is a G protein-coupled receptor (GPCR) that senses extracellular hydrogen ions. Stimulation of TDAG8 elicits a signaling cascade that leads to the phosphorylation of cAMP-responsive element-binding protein 1 (CREB1). Activated CREB1 is translocated to the nucleus, where it binds to a synthetic promoter (P<sub>CRE</sub>) containing cAMP-response elements (CRE) and serves as a transcription factor. Thus, the authors genetically linked the increase in hydrogen ion concentration (a biomarker of KDA) to secretion of insulin in HEK<sub>pH-Guard</sub> cells (Figure 3D).

Obesity is a common metabolic disorder (Chooi et al., 2019), and is associated with the development of cardiovascular disease, diabetes, and cancer (Blüher, 2019). Although obesity can be prevented or reversed by lifestyle changes, such as restriction of dietary intake and physical exercise, many obese patients still require medical treatment. Currently, the main treatment options are anti-obesity medications, which are generally only partially effective, or bariatric surgery (Ruban et al., 2019), which can lead to serious and sometimes fatal complications. Rössger et al. engineered designer cells that can ameliorate diet-induced obesity in mice (Rössger et al., 2013). They utilized HEK-293T cells in which stable expression of a synthetic free fatty acid-induced transcription factor is biochemically connected to the secretion of pramlintide, a clinically licensed appetite-suppressing peptide hormone (Figure 3E). The engineered transcription factor is a chimeric protein consisting of a phloretin-responsive repressor (TtgR) fused to peroxisome proliferator-activated receptor-α (PPARα). Exposure of the cells to high blood levels of free fatty acids releases the inhibitory effects of various endogenous repressors on the PPARα chimera, leading to expression of pramlintide via a TtgR-specific promoter.

Elevated uric acid levels, known as hyperuricemia, cause multiple metabolic disorders such as gout and tumor lysis syndrome (Skoczyńska et al., 2020). Uric acid, a by-product of purine metabolism, is barely soluble
in aqueous environments, and high levels of this compound tend to deposit as crystals in joints, kidneys, and other tissues in the body. Kemmer et al. constructed a synthetic, cell-based system based on a modified Deinococcus radiodurans-derived protein that senses uric acid levels and elicits the secretion of engineered Aspergillus flavus urate oxidase, which promotes uric acid excretion (Kemmer et al., 2010). This closed-loop system incorporates a bacterial transcriptional repressor (HucR) with high affinity for a DNA sequence motif (hucO) in a uric acid-free environment (Figure 3F). In the presence of uric acid at a sufficient concentration, HucR dissociates from DNA, leading to initiation of gene expression. To further enhance the repression, HucR was C-terminally fused to a Krueppel-associated box (KRAB) domain. This device successfully restored the urate concentration to normal levels in urate oxidase-deficient mice.

Graves’ disease is an autoimmune disorder caused by the generation of autoantibodies that stimulate thyroid-stimulating hormone (TSH) receptors in the thyroid gland, leading to constitutive and chronic over-secretion of these hormones (T3 and T4) into the blood (Davies et al., 2020). Current therapeutic interventions include antithyroid medications, radioactive iodine treatment, and in some cases thyroidectomy followed by administration of exogenous thyroid hormones. Saxena et al. engineered designer cells equipped with a self-regulating sensor–effector TSR-TSHAntag circuit (Saxena et al., 2016). This system contains a chimeric protein consisting of the ligand-binding domain of TR fused with Gal-4, a yeast DNA-binding domain (Figure 3G). In the presence of high circulating levels of thyroid hormones T3 and T4, this hybrid protein binds to Ga-l4 response element and initiates a complex transcription process involving steroid receptor coactivator-1 (SRC-1) and TRAP 220, thereby promoting histone acetylation and resulting in gene expression. As a therapeutic output, they employed thyroid-stimulating hormone receptor antagonist (TSHAntag), which competitively inhibits TSHR in the thyroid gland, thus decreasing the production of thyroid hormone T3 and T4 and alleviating the disease state.

DESIGNER CELLS WITH ANTIMICROBIAL FUNCTIONALITY

It has been estimated that more than 15 million people die each year from infectious diseases or their complications (Ahrens and Pigeot, 2014). Currently, the main treatment for bacterial infections is antibiotics, but drug resistance is becoming an ever-increasing problem (Aslam et al., 2018). Recently, synthetic biologists have started to develop engineered mammalian cells designed specifically to eliminate multidrug-resistant (MDR) organisms. Liu and coworkers focused on designer cells with the ability to efficiently detect and eradicate meticillin-resistant Staphylococcus aureus (MRSA) – a difficult-to-treat superbug (Liu et al., 2018). They engineered HEK-293T cells expressing toll-like-receptors (TLRs) that recognize various epitopes presented on the bacterial membrane. When these immunomimetic cells encounter and bind to bacteria, they are triggered to secrete lysostaphin, a bacteriolytic enzyme highly lethal to S. aureus (Figure 4A). The binding of TLRs to bacterial membrane-anchored molecules activates NF-kB signaling, resulting in the expression of a therapeutic gene, in this case lysostaphin, under an NF-kB-responsive promoter. These designer cells were highly effective in treating MRSA infections both in vitro and in a mouse model.

Another strategy is to focus on biofilm disintegration. Biofilm is a complex structure composed of aggregates of bacterial cells interconnected by endogenously produced polymers (Flemming et al., 2016). Bacterial strains that form biofilms are generally hard to treat, since they can physically evade the immune system because of the cross-linked jelly-like environment of the biofilm, which also restricts the access of antibiotics. The consequent continuous exposure to subtherapeutic levels of antibiotics further increases the resistance of the bacteria. Thus, targeting the biofilm matrix seems a promising approach to treat MDR strains. Sedlmayer et al. engineered nonimmune designer cells designed to selectively block Pseudomonas aeruginosa biofilm formation (Sedlmayer et al., 2017). They provided HEK-293T cells with a synthetic gene network harboring a PAI-1-activated transcription factor. PAI-1 is a bacterial autoinducer secreted by Pseudomonas aeruginosa that is involved in cellular communication (quorum sensing); it assists the bacteria to evade the immune response (Figure 4B). This sensor-transcription hybrid consists of P. aeruginosa LasR-derived activator fused to VP-16 transcriptional transactivator. The exposure of these quorum-quencher cells to P. aeruginosa PAI-1 triggers transcription of genes controlled by a LasR-response element. They utilized three biofilm-dissolving outputs, glycoside hydrolases PsIgH and PeIAH, and autoinducer-inactivating lactonase (MomL), with the aim of achieving synergistic action. The results suggest that this is a promising approach for the treatment of infections involving biofilm-forming bacteria.

DESIGNER CELLS TO TREAT INFLAMMATORY DISEASES

Cytokines are peptide-based signaling molecules that play a major role in cellular communication and the immune response (Turner et al., 2014). Abnormal cytokine secretion contributes to many conditions, such as
inflammatory and autoimmune diseases, cancer, and neurological disorders (Kany et al., 2019). For example, psoriasis is a chronic inflammatory skin disease characterized by the overproduction of pro-inflammatory cytokines, particularly tumor necrosis factor (TNF) and interleukin-2 (IL22) (Rendon and Schäkel, 2019). Because there is no cure, the main therapeutic approaches focus on symptomatic treatment, including the use of antibodies that neutralize pro-inflammatory cytokines and immunosuppressant drugs such as methotrexate and cyclosporine. Though these agents provide symptomatic relief, they are associated with severe side effects, such as recurrent infections and immunogenicity. Recent clinical trials have demonstrated that the administration of immunomodulatory cytokines IL4 (22) and/or IL10 (23) at well-tolerated doses significantly improves psoriasis. Nevertheless, these interleukins are quickly eliminated from the blood circulation, necessitating frequent injections. Schukur et al. developed cytokine-converter designer cells for autonomous management of psoriasis (Schukur et al., 2015). In their system, an AND-logic gate was designed using TNFα and IL22 as signal inputs, and IL4 plus IL10 as a secreted output (Figure 5A). They engineered HEK-293T cells with TNFα receptor (TNFR) rewired to the expression of the bactericidal protein lysostaphin.

DESIGNER CELLS FOR CANCER TREATMENT
The well-established chimeric antigen receptor (CAR)-T technology is a cell-based therapy that is clinically effective against certain types of malignancies, with long-lasting beneficial effects (June et al., 2018).
Autologous T-cells are isolated from patients and then genetically engineered to express synthetic CARs, which function to redirect T cells to selectively recognize and destroy cancerous cells bearing specified epitopes on their surface (Figure 6A). The FDA has approved the use of CAR-T cells to treat hematopoietic cancers. However, this system still has serious drawbacks, which limit its clinical use. These include life-threatening adverse effects and cell-associated toxicities, inadequate efficacy against solid tumors, the development of resistance, and antigen escape (Sterner and Sterner, 2021).

Figure 5. Designer cells for the management of psoriasis

(A) The design of autonomous therapeutic cells with an AND-ON gate for psoriasis treatment. Cells were equipped with TNF receptor (TNFR), NFκB-regulated human IL22 receptor alpha unit (hIL22R), and stat3-driven mouse IL4 and IL10 (mIL4 and mIL10) genes. In psoriatic patients, high levels of TNF bind to its receptor and activate the expression of hIL22RA alpha via NFκB. The newly synthesized hIL22RA is translocated to the membrane where it forms a functional heterodimer with IL10RB, initiating a signaling cascade that results in the phosphorylation of STAT3. The phosphorylated STAT3 forms a functional dimer that is translocated to the nucleus, resulting in enhanced expression of anti-psoriatic interleukins, mIL4 and mIL10.

CONCLUSION AND OUTLOOK

Designer cell-based therapeutic technology has already demonstrated its superiority over conventional drugs for the management of some difficult-to-treat diseases. An example of their clinical potential is provided by CAR-T cells, which are already in clinical use to cure various life-threatening hematopoietic tumors. Furthermore, autonomously operating designer cells would likely improve compliance and quality of life of patients with chronic diseases requiring invasive treatment, such as diabetes. However, a number of challenges remain.
The possibility of off-target induction (unexpected and undesired activation) of designer cells should be taken into account during their engineering for therapeutic purposes. The nature of the signaling circuit that connects the input-sensing transducer to effector modules has a significant impact on the performance and safety profile of designer cells. A totally orthogonal signaling cascade is generally required to build safe and reliable designer cells in order to rule out undesired induction. Orthogonal signaling refers to an ectopically introduced signaling network that is not influenced by endogenous pathways of the organism, thus ensuring tightly controlled and selective activation. Because many genetically encoded transducers are naturally wired to endogenous pathways, engineering of novel sensing transducers with the ability to orthogonally stimulate designer cells will be an important avenue for future research. Alternatively, engineering of designer cells with an AND-gate operation mode, that is, an activation mechanism which functions only in simultaneous presence of at least two different inducers, can be used to reduce off-target induction. However, the engineering of such logic gates is not an easy task, as it requires the availability of at least two different, reliable biomarkers for the same disease.

Open-loop operating expression systems can offer ON/OFF behavior, depending on the inducer availability in the cell environment. Although open-loop-based devices required repetitive administration of the inducer, a combination of open and closed loops may be very highly tunable, as well as potentially having the ability to temporarily block designer cell function if necessary or desirable. For example, if autonomous designer cells could be turned off by an external inducer, this would provide another control point in the event of therapeutic-induced toxicity or if the cell implant does not function as intended. Designer cells equipped with such an “emergency stop button” would offer a new layer of safety, especially for cell implants intended for long-term therapeutic use.

In fact, typical biomarkers are not available for all diseases, making the design of closed-loop gene circuits problematic. It has recently been demonstrated that specific extracellular RNAs (exRNAs) molecules are secreted into the body circulation in certain disease states (Xi et al., 2017). For example, several studies have characterized a number of distinctive exRNA molecules that are associated with Alzheimer’s disease, cancers, and numerous inflammatory diseases (Roser et al., 2018; Xu et al., 2021). Thus, the bioengineering of selective and sensitive cell-based molecular architectures that can detect and utilize these exRNAs as...
signal inputs would significantly increase the disease biomarker repertoire, facilitating the design of therapeu tic designer cells for a wide range of diseases. Despite all the challenges that are still associated with their engineering, it is already clear that therapeutic designer cells will play a pivotal role in next-generation medicine.

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DECLARATION OF INTERESTS

The authors declare no competing interests.

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