An engineered \(\text{L-arginine}\) sensor of *Chlamydia pneumoniae* enables arginine-adjustable transcription control in mammalian cells and mice

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**ABSTRACT**

For optimal compatibility with biopharmaceutical manufacturing and gene therapy, heterologous transgene control systems must be responsive to side-effect-free physiologic inducer molecules. The arginine-inducible interaction of the ArgR repressor and the ArgR-specific ARG box, which synchronize arginine import and synthesis in the intracellular human pathogen *Chlamydia pneumoniae*, was engineered for arginine-regulated transgene (ART) expression in mammalian cells. A synthetic arginine-responsive transactivator (ARG), consisting of ArgR fused to the *Herpes simplex* VP16 transactivation domain, reversibly adjusted transgene transcription of chimeric ARG box-containing mammalian minimal promoters (PART) in an arginine-inducible manner. Arginine-controlled transgene expression showed rapid induction kinetics in a variety of mammalian cell lines and was adjustable and reversible at concentrations which were compatible with host cell physiology. ART variants containing different transactivation domains, variable spacing between ARG box and minimal promoter and several tandem ARG boxes showed modified regulation performance tailored for specific expression scenarios and cell types. Mice implanted with micro-encapsulated cells engineered for ART-inducible expression of the human placental secreted alkaline phosphatase (SEAP) exhibited adjustable serum phosphatase levels after treatment with different arginine doses. Using a physiologic inducer, such as the amino acid \(\text{L-arginine}\), to control heterologous transgenes in a seamless manner which is devoid of noticeable metabolic interference will foster novel opportunities for precise expression dosing in future gene therapy scenarios as well as the manufacturing of difficult-to-produce protein pharmaceuticals.

**INTRODUCTION**

Conditional transcription control systems which fine-tune heterologous transgene expression in mammalian cells are fundamental for gene-function analysis (1,2), drug discovery (3,4), expression dosing in gene therapy (5), design of synthetic gene networks and for manufacturing of difficult-to-produce protein pharmaceuticals (6,7). Currently available mammalian transgene control systems capitalize on a generic design consisting of a synthetic transactivator (transrepressor), typically derived from a prokaryotic response regulator fused to a mammalian transactivation (transrepression) domain and a chimeric promoter assembled by cloning transactivator-specific operator sites adjacent to a minimal (constitutive) eukaryotic promoter (8–10). Binding of an inducer molecule modulates the affinity of the transactivator (transrepressor) for the cognate promoter and either induces (ON-type systems, (11–14)) or represses (OFF-type systems, (14,15)) transcription of linked transgenes. Trigger molecules and parameters include antibiotics (10,14,16), hormones and hormone analogs (17–19), quorum sensing substances (12,20,21), temperature (22), the redox poise (20), immunosuppressive and antidiabetic drugs (23,24), gaseous acetaldehyde (11) and biotin (25). Although most of these established gene regulation systems show excellent regulation performance *in vitro* as well as in animals, the inducer molecules are often incompatible with gene therapy and biopharmaceutical manufacturing scenarios because of their side-effects, which are well documented for hormones and some of their analogs (26,27), antibiotics (28–30) and immunosuppressive drugs (31,32). While the biopharmaceutical manufacturing industry strives to develop gene regulation systems controlled by FDA-approved media components...
to prevent prohibitive downstream processing and lic-
ing procedures (33), the gene therapy community dreams of capturing pathologic signals which can be converted into a well-tuned therapeutic intervention. The common interest is exploiting physiologic signal molecules for titration of heterologous product gene expression and prevent interference between endogenous and synthetic regulatory circuits.

As an obligate intracellular pathogen infecting the respiratory tracts and causing pneumonia and athero-
sclerotic heart disease (34–37), *Chlamydia pneumoniae* has evolved to capture physiologic signals and synchronize its metabolism with the host cell (38). For example, the coordination of arginine biosynthesis and catabolism with availability of this amino acid in host cells is managed by ArgR, a master regulatory molecule acting as an arginine-dependent apo-repressor that specifically binds to arginine-responsive transactivator (ARG) box operators and represses the *glnPQ* operon which encodes a putative arginine transport system in an arginine-responsive manner (39,40). ArgR, which is highly conserved in bacterial systems, can function both as transcriptional repressor or activator (41,42). In *Escherichia coli*, ArgR negatively regulates the expression of the arginine biosynthetic genes in response to intracellular \( \mathcal{L} \)-arginine levels (43). When acting as a repressor, ArgR requires allosteric activation by \( \mathcal{L} \)-arginine to bind to an 18-bp palindromic operator sequence known as the ARG box (39,43–45). While the N-terminal half of ArgR contains a winged helix-turn-helix family DNA binding domain, the C-terminal half is responsible for arginine binding.

We have engineered ArgR and the ARG box of *C. pneumoniae* for *\( \mathcal{L} \)*-arginine-regulated transgene (ART) regulation system in mammalian cells. As an ON-type system by default, ART remained silent at minimal endogenous *\( \mathcal{L} \)*-arginine concentrations and was adjustably induced by increasing *\( \mathcal{L} \)*-arginine levels. Using a proteogenic amino acid to control heterologous transgene in mammalian cells grown in monolayer cultures or in mice should foster new opportunities in gene therapy and biopharmaceutical manufacturing.

**MATERIALS AND METHODS**

**Plasmid construction**

All plasmids and oligonucleotides constructed and used in this study are listed in Table 1. Details on vector design are also provided in Table 1.

**Cell culture, transfection and construction of stable cell lines**

Wild-type Chinese hamster ovary cells (CHO-K1, ATCC: CCL-61) and its derivatives [e.g. CHO-ET, SEAP], (10)] were cultivated in *\( \mathcal{L} \)*-arginine-free Hektor G (Cell Culture Technologies, Gravesano, Switzerland) supplemented with 10 mg/l *\( \mathcal{L} \)-arginine, 5% (v/v) fetal calf serum (FCS, PAN Biotech GmbH, Aidenbach, Germany, Cat. No. 3302, Lot No. P251110) and 1% (v/v) penicillin/streptomycin solution (Sigma, St Louis, MO, USA, Cat. No. P4458). Human fibrosarcoma cells (HT-1080, ATCC, CCL-121), African green monkey kidney cells (COS-7, ATCC CRL-1651) and human embryonic kidney cells, transgenic for the simian virus 40 (SV40) large T antigen [HEK293-T, (46)], were cultivated in *\( \mathcal{L} \)*-arginine-free Hektor G (Cell Culture Technologies, Gravesano, Switzerland) supplemented with 10 mg/l *\( \mathcal{L} \)-arginine, 10% (v/v) FCS and 1% (v/v) penicillin/streptomycin solution. Mouse fibroblasts (NIH/3T3, ATCC: CRL-1658) were cultured in Dulbecco's modified Eagle's medium (DMEM, Invitrogen, Basel, Switzerland, Cat. No. 52100-39) supplemented with 10% FCS and 1% (v/v) penicillin/streptomycin solution. All cell types were cultivated at 37°C in a humidified atmosphere containing 5% CO₂.

For DNA transfection, 1.2 μg DNA (for co-transfection an equal amount of each plasmid) was transfected into 55000 cells pre-cultivated for 24 h in a well of a 24-well plate. For transfection of CHO-K1, the plasmid DNA was diluted into a total volume of 25 μl 0.5 M CaCl₂ solution, mixed with 25 μl PO₄ solution (100 mM HEPES, 280 mM NaCl, 1.5 mM Na₂HPO₄, pH 7.1) and incubated for 15 min at room temperature. The DNA–calcium phosphate complex was added, and centrifuged onto the cells (5 min at 12000 g) to increase transfection efficiency. After incubation for 2 h, the cells were exposed to a glycerol shock for 30 s (*\( \mathcal{L} \)-arginine-free ChoMaster® HTS medium supplemented with 15% glycerol). After a single washing with phosphate-buffered saline (PBS, Dulbecco's phosphate-buffered saline, Invitrogen, Basel, Switzerland, Cat. No. 21600-0069), the cells were cultivated in ChoMaster® HTS supplemented with different *\( \mathcal{L} \)-arginine concentrations. For transfection of COS-7, HEK293-T, the standard calcium phosphate-based transfection protocol was used (21) and transfected populations were cultivated in Hektor G supplemented with different *\( \mathcal{L} \)-arginine concentrations. HT-1080 and NIH/3T3 were transfected with Fugene™ 6 (Roche Diagnostics AG, Basel, Switzerland, Cat. No. 11814443001) following the supplier's protocol and cultivated in Hektor G and DMEM, respectively. Production of the reporter protein was assessed in the culture supernatants 60 h after transfection, unless indicated otherwise.

CHO-ARG2-SEAP, transgenic for *\( \mathcal{L} \)-arginine-controlled SEAP expression, was constructed by sequential co-transfection and clonal selection of (i) pSH91 and pPur (Clontech, Cat. No. 6156-1) (CHO-ARG2) and (ii) pSH93 and pSV2neo (Clontech, Cat. No. 6172-1) (CHO-ARG2-SEAP). To assess the dose–response characteristics of ART-regulated gene expression, CHO-ARG2-SEAP was cultured at 110000 cells/ml for 60 h in *\( \mathcal{L} \)-arginine-free* ChoMaster® HTS supplemented with *\( \mathcal{L} \)-arginine ranging from 10 mg/l to 10000 mg/l. Reversibility of *\( \mathcal{L} \)-arginine-mediated SEAP production was assessed by cultivating CHO-ARG2-SEAP (110000 cells/ml) for 1 week while alternating *\( \mathcal{L} \)-arginine concentrations from 10 to 1000 mg/l every 48 h.

**Western blot analysis**

Sixty hours after transfection of 350 000 HEK-293T cells with the vector encoding ARG2 (pSH91, PSV40-ARG2-pA), the cell lysate was heat denaturated in
Table 1. Plasmids and oligonucleotides used and designed in this study

| Plasmid | Description and cloning strategy | Reference or source |
|---------|----------------------------------|---------------------|
| pRevTRE | Oncoretroviral expression vector containing a tetracycline-responsive expression unit | Clontech, Palo Alto, CA, USA |
| pPur | Selection vector conferring puromycin resistance to eukaryotic cells | Clontech, Palo Alto, CA, USA |
| pSV2neo | Selection vector conferring neomycin resistance to eukaryotic cells | Clontech, Palo Alto, CA, USA |
| pBP10 | Vector encoding a P ETR5-driven SEAP expression unit (P ETR5-SEAP-pA; P ETR5-2bp-P CMVmin) | (9) |
| pBP11 | Vector encoding a P ETR5-driven SEAP expression unit (P ETR5-SEAP-pA; P ETR5-4bp-P CMVmin) | (9) |
| pBP12 | Vector encoding a P ETR7-driven SEAP expression unit (P ETR7-SEAP-pA; P ETR7-6bp-P CMVmin) | (9) |
| pBP13 | Vector encoding a P ETR7-driven SEAP expression unit (P ETR7-SEAP-pA; P ETR7-8bp-P CMVmin) | (9) |
| pBP14 | Vector encoding a P ETR9-driven SEAP expression unit (P ETR9-SEAP-pA; P ETR9-10bp-P CMVmin) | (9) |
| pMF111 | Vector encoding a P CMVmin-driven SEAP expression unit (P CMVmin-SEAP-pA) | (69) |
| pMT1227 | Vector encoding the C. pneumoniae CWL029 ArgR gene | (39) |
| pWW35 | Constitutive ET1 expression vector (P SV40-ET1-pA) | (10) |
| pWW42 | Constitutive ET2 expression vector (P SV40-ET2-pA) | (10) |
| pWW64 | Constitutive ET3 expression vector (P SV40-ET3-pA) | (9) |
| pSH91 | Constitutive ARG2 expression vector (P SV40-ARG2-pA) | This work |
| pSH92 | Vector encoding a P ART1-driven expression unit (P ART1-SEAP-pA; P ART1-O ARG-0bp-PhCMVmin) | This work |
| pSH93 | Vector encoding a P ART2-driven expression unit (P ART2-SEAP-pA; P ART2-O ARG-2bp-P CMVmin) | This work |
| pSH94 | Vector encoding a P ART3-driven expression unit (P ART3-SEAP-pA; P ART3-O ARG-4bp-P CMVmin) | This work |
| pSH95 | Vector encoding a P ART4-driven expression unit (P ART4-SEAP-pA; P ART4-O ARG-6bp-P CMVmin) | This work |
| pSH96 | Vector encoding a P ART5-driven expression unit (P ART5-SEAP-pA; P ART5-O ARG-8bp-P CMVmin) | This work |
| pSH97 | Vector encoding a P ART6-driven expression unit (P ART6-SEAP-pA; P ART6-O ARG-10bp-P CMVmin) | This work |
| pSH98 | Vector encoding a P ARTm1-driven expression unit (P ARTm1-SEAP-pA; P ARTm1-O ARG-0bp-P CMVmin) | This work |
| pSH99 | Vector encoding a P ARTm2-driven expression unit (P ARTm2-SEAP-pA; P ARTm2-O ARG-2bp-P CMVmin) | This work |
| pSH100 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-4bp-P CMVmin) | This work |
| pSH101 | Vector encoding a P ARTm4-driven expression unit (P ARTm4-SEAP-pA; P ARTm4-O ARG-6bp-P CMVmin) | This work |
| pSH102 | Vector encoding a P ART5-driven expression unit (P ART5-SEAP-pA; P ART5-O ARG-8bp-P CMVmin) | This work |
| pSH103 | Vector encoding a P ART6-driven expression unit (P ART6-SEAP-pA; P ART6-O ARG-10bp-P CMVmin) | This work |
| pSH104 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH105 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH106 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH107 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH108 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH109 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH110 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH111 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH112 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH113 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH114 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH115 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH116 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH117 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH118 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH119 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH120 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH121 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |
| pSH122 | Vector encoding a P ARTm3-driven expression unit (P ARTm3-SEAP-pA; P ARTm3-O ARG-7bp-P CMVmin) | This work |

(Continued)
Table 1. Continued

| Plasmid    | Description and cloning strategy | Reference or source         |
|------------|----------------------------------|----------------------------|
| pSH127     | Vector encoding a P_{ART\text{m4}}-driven expression unit (P_{ART\text{m4}}-SEAP-pA; P_{ART\text{m4}}-Ascl-OARG-7bp-OARG-7bp-OARG-MluI-0bp-P_{hCMVmin}) | This work                   |
| P_{hCMVmin}| P_{hCMVmin} minimal version of the human cytomegalovirus promoter |                            |
| P_{SV40}   | P_{SV40} constitutive simian virus 40 promoter |                            |
| SEAP       | SEAP human placental secreted alkaline phosphatase |                            |
| VP16       | human E2F4 transactivation domain |                            |
| ET1        | macrolide-dependent transactivator (E-VP16) |                            |
| ET2        | macrolide-dependent transactivator (E-p65) |                            |
| ET3        | macrolide-dependent transactivator (E-E2F4) |                            |
| OARG-7bp-OARG–7bp-OARG-7bp-OARG-MluI-0bp-P_{hCMVmin} |                            |
| ArgR, transactivator of the C. pneumoniae CWL029 of the glnPQ operon |                            |
| ARG1       | L-arginine-dependent transactivator (ArgR-p65); ArgR, transactivator of the early promoter |                            |
| ARG2       | L-arginine-dependent transactivator (ArgR-VP16); ArgR3, L-arginine-dependent transactivator (ArgR-E2F4); E2F4, transactivation domain of the human E2F4 |                            |
| ARG3       | E2F4, transactivation domain of the human E2F4; ET1, macrolide-dependent transactivator (E-VP16); ET2, macrolide-dependent transactivator (E-p65); ET3, macrolide-dependent transactivator (E-E2F4); ETR, operator module specific for MphR(A); IRES_{PU}, internal ribosome entry site of polioviral origin |                            |
| P_{hCMVmin}| P_{hCMVmin} minimal version of the human cytomegalovirus promoter |                            |
| P_{SV40}   | P_{SV40} constitutive simian virus 40 promoter |                            |

Regulating l-arginine and l-arginine analogs

L-arginine stock solution was prepared in water by adjusting 50 g/l L-arginine base (0.287 M, AppliChem, Darmstadt, Germany, Cat. No. A36530100) to pH 7.2 with HCl and used at the final concentrations indicated. L-ornithine hydrochloride (Fluka, Buchs, Switzerland, Cat. No. 75470) and l-citrulline (Fluka, Buchs, Switzerland, Cat. No. 27510), l-homoarginine hydrochloride (Acros Organics, Basel, Switzerland, Cat. No. 1690900050) and L-arginine ethyl ester dihydrochloride (Fluka, Buchs, Switzerland, Cat. No. 11030), L-arginine ethyl ester dihydrochloride (Sigma-Aldrich, Steinheim, Germany, Cat. No. A2883), agmatine sulfate (Sigma-Aldrich, Cat. No. A7127) and L-canavanine sulfate (Sigma-Aldrich, Cat. No. C9758) stock solutions were prepared by adjusting the pH of a 0.144 M solution to pH 7.2.

Quantification of reporter gene expression

Production of the human placental secreted alkaline phosphatase (SEAP) was quantified using a p-nitrophenylphosphate-based light absorbance time course (47,48). Interferon-β quantification was determined using the human Interferon β-specific ELISA (PBL Laboratories, NJ, USA, Cat. No. 41400-1) according to the manufacturer’s protocol.

Determination of osmolality

The osmolality of ChoMaster® HTS medium supplemented with different L-arginine concentrations was assessed with a Vapro® according to the manufacturer’s instructions (Wescor Inc., Logan, UT, USA).

In vivo methods

CHO-K1 cells, engineered for L-arginine-controlled SEAP expression (CHO-ARG2-SEAP), were encapsulated into 400 μm alginate-poly-(l-lysine)-alginate beads (200 cells/capsule) using the Inotech Encapsulator Research IE-50R (Inotech Biotechnologies Ltd, Basel, Switzerland) according to the manufacturer’s protocol and the following parameters: 0.2 mm nozzle, 20 ml syringe at a flow rate of 405 units, nozzle vibration frequency 1024 Hz, voltage for bead dispersion 900 V. Seven hundred microliters of l-arginine-free ChoMaster® HTS containing 2 × 10^6 encapsulated cells (10^6 capsules/mouse) were injected intraperitoneally into mice (oncins France souche 1, Charles River Laboratories, France). One hour after implantation, l-arginine was administered by injection at doses ranging from 0 to 100 mg/kg. l-arginine was prepared for in vivo administration by diluting the l-arginine stock solution (see above) with an iso-osmotic 0.9% NaCl solution to obtain the appropriate concentrations. Control mice were implanted with capsules containing wild-type CHO-K1 cells. Seventy-two hours after implantation, the mice were sacrificed and their blood collected. For SEAP quantification, serum was isolated using a microtainer SST tube (Beckton Dickinson, Plymouth, UK) according to the manufacturer’s protocol. All experiments involving mice were performed according to the European Community Council directive (86/609/ EEC), approved by the French Ministry of Agriculture and Fishery (Paris, France) and performed by M.D.E.-B. at the Institut Universitaire de Technologie, IUTA, F-69622 Villeurbanne Cedex, France.
RESULTS

Determination of the L-arginine concentration range compatible with host physiology and potential L-arginine-controlled gene expression

Since L-arginine plays a central role in the urea cycle, the synthesis of nitric oxide and for the synthesis of cytoplasmic and nuclear proteins (49), it is often the rate-limiting amino acid in fast-growing organisms (50,51). In order to prevent L-arginine becoming a physiologic bottleneck or endogenous L-arginine interfering with ART-controlled transgene expression in mammalian cells, we exposed a CHO-K1-derived SEAP-expressing cell line [CHO-ET1-SEAP1, (10)] to L-arginine concentrations ranging from 0 to 10 000 mg/l and scored SEAP levels and cell density after a 60 h cultivation period. A 10 mg/l L-arginine was sufficient to support robust wild-type-like cell growth with SEAP production levels equivalent to control cultivations in standard ChoMaster® HTS medium containing a default L-arginine concentration of 200 mg/l (Figure 1). This observation is quantitatively comparable with physiologic plasma concentrations of L-arginine found in mice and humans (~17 mg/l (52)). Between 10 mg/l and 2 g/l of L-arginine, SEAP production and cell densities were comparable to populations cultivated in standard ChoMaster® HTS medium. At L-arginine concentrations above 2 g/l the culture media became hyperosmotic (>310 mOsm) which compromised cell viability and titer (Figure 1). Based on this data, we selected 10 mg/l and 1 g/l as low and high L-arginine concentrations in follow-up experiments.

Design of a L-arginine-responsive mammalian transcription control system

Capitalizing on the ArgR repressor of *Chlamydia pneumoniae*, which manages arginine metabolism by binding to specific operator sequences (O<sub>ARG</sub>) upstream of the *glnPQ* operon in a L-arginine-inducible manner, we have designed a L-arginine-regulated transgene control system (ART) which fine-tunes transgene transcription in mammalian cells. ART consists of two components, a L-arginine-regulated transactivator (ARG1) and an ARG1-specific synthetic promoter (P<sub>ART</sub>): (i) ARG1 was designed by fusing ArgR C-terminally to the human NF-kB transactivation domain (ARG1, ArgR-p65; pSH120, P<sub>Sva40-ARG1-pA</sub>) (8,39). (ii) P<sub>ART</sub> was assembled by cloning an ArgR-specific operator module harboring two ARG boxes separated by a single base pair triplet (O<sub>ARG</sub>, AGTTTCTTGGATTAATTGG (predicted ARG boxes underlined)) 5’ of a minimal version of the human cytomegalovirus immediate early promoter (P<sub>CMVmin</sub>; P<sub>ART</sub>, O<sub>ARG</sub>-P<sub>CMVmin</sub>; pSH93, P<sub>ART</sub>-SEAP-pA; Figure 2A) (15,39). Upon co-transfection of pSH120 (P<sub>Sva40-ARG1-pA</sub>) and pSH93 (P<sub>ART</sub>-SEAP-pA) into CHO-K1, SEAP production was profiled after cultivation for 60 h in medium containing either 10 mg/l or 1 g/l L-arginine (pSH120 and pSH93: 1.12 ± 0.07 U/l (10 mg/l L-arginine), 7.25 ± 0.39 U/l (1 g/l L-arginine); pSH91 and pSH93: 0.50 ± 0.07 U/l (10 mg/l L-arginine), 10.66 ± 0.09 U/l (1 g/l L-arginine); pSH121 and pSH93: 0.75 ± 0.22 U/l (10 mg/l L-arginine), 2.76 ± 0.08 U/l (1 g/l L-arginine). SEAP expression values of the different transactivator–promoter combinations exhibited varied performance characteristics with the basal expression of ARG2 being slightly lower compared to ARG1 and ARG3 exhibiting the poorest performance characterized by lower maximum and higher basal expression levels. Taken together, co-transfection of P<sub>ART</sub> with ARG2 enabled the highest induction performance for SEAP expression (Figure 3B–D, SEAP values with P<sub>ART</sub>). The relative spacing of promoter components such as operator and minimal promoter influences the assembly of operon by ArgR in *Chlamydia pneumoniae* (39) (Figure 2B and C).

Optimizing the ART system I—different L-arginine-dependent transactivator variants

Weber and colleagues (9) have demonstrated that the type of transactivation domain fused to the response regulator may impact overall regulation performance. To characterize potential alternatives to the p65 transactivation domain, we fused the *Chlamydia pneumoniae* L-arginine-dependent transactivator to the VP16 domain from *Herpes simplex* (ARG2) (53) and the transactivation domain of the human E2F4 (ARG3) (54) (Figure 3A). For comparative analysis of the regulation performance, ARG1-, ARG2- and ARG3-encoding plasmids were co-transfected with pSH93 (P<sub>ART</sub>-SEAP-pA) into CHO-K1 and SEAP production was profiled after cultivation for 60 h in medium containing either 10 mg/l or 1 g/l L-arginine (P<sub>SH120</sub> and pSH93: 1.12 ± 0.12 U/l (10 mg/l L-arginine), 0.07 U/l (10 mg/l L-arginine), 0.22 U/l (1 g/l L-arginine), 2.76 ± 0.08 U/l (1 g/l L-arginine). SEAP expression values of the different transactivator–promoter combinations exhibited varied performance characteristics with the basal expression of ARG2 being slightly lower compared to ARG1 and ARG3 exhibiting the poorest performance characterized by lower maximum and higher basal expression levels. Taken together, co-transfection of P<sub>ART</sub> with ARG2 enabled the highest induction performance for SEAP expression (Figure 3B–D, SEAP values with P<sub>ART</sub>).
The L-arginine-responsive promoter (P\text{ART1}) harbors an ArgR-specific operator sequence (O\text{ARG}, capital letters, ARG boxes underlined) and drives expression of a reporter gene (e.g. human secreted alkaline phosphatase, SEAP) in a L-arginine-dependent manner under the control of the simian virus 40 promoter (P\text{SV40}). Transactivation domain (ARG1, ArgR-p65), is expressed in a constitutive manner upon binding to P\text{ART1} through direct ARG1-O\text{ARG} interaction, thus enabling the transcription of the reporter gene.

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Increasing the number of tandem operator modules within a trigger-inducible promoter may increase maximum expression levels as more transactivators can be recruited to the promoter. The transcription activity of the promoters harboring the operator in a minimal expression levels (10 mg/l L-arginine) (Figure 3C). Overall, P\text{ART1} showed an optimal balance between basal and maximum expression.

**Optimizing the ART system III—promoter variants which differ in the number of ARG box operator modules**

Increasing the number of tandem operator modules within a trigger-inducible promoter may increase maximum expression levels as more transactivators can be recruited to the promoter. The transcription activity of the promoters harboring the operator in a minimal expression levels (10 mg/l L-arginine) (Figure 3C). Overall, P\text{ART1} showed an optimal balance between basal and maximum expression.

The bacterial repressor ArgR of *Chlamydia pneumoniae*, fused to the human NF-κB transactivation domain (ARG1, ArgR-p65), is expressed in a constitutive manner under the control of the simian virus 40 promoter (P\text{SV40}). The L-arginine-responsive promoter (P\text{ART1}) harbors an ArgR-specific operator sequence (O\text{ARG}, capital letters, ARG boxes underlined) upstream of the minimal version of the cytomegalovirus immediate early promoter (P\text{hCMVmin}) and drives expression of a reporter gene (e.g. human secreted alkaline phosphatase, SEAP) in a L-arginine-induced manner. All expression units are terminated by a polyadenylation site (pA). Selected restriction sites are indicated: A, NcoI; B, BssHII; Ba, BamHI; E, EcoRI; H, HindIII; N, NotI; S, SfiI; X, XhoI, Xh, XhoI. (B) At a low L-arginine concentration (10 mg/l), the ARG1 is in a low-affinity DNA binding state and does not interact with its specific operator sequence (O\text{ARG}); therefore, expression of the transgene remains silent. At a higher L-arginine concentration (1 g/l), the ARG1 is in a high-affinity DNA binding state and interacts with its specific operator sequence (O\text{ARG}) and activates transcription from P\text{ART1}.

For co-transfection of CHO-K1 with pSH120, E2F4 transactivation domain mediated the poorest regulation performance, characterized by higher leakiness (10 mg/l L-arginine) and lower maximum expression levels (1 g/l L-arginine) (Figure 3C). Overall, P\text{ART1} showed an optimal balance between basal and maximum expression.

**Figure 2.** Diagram of the ART regulation system. (A) The bacterial repressor ArgR of *Chlamydia pneumoniae*, fused to the human NF-κB transactivation domain (ARG1, ArgR-p65), is expressed in a constitutive manner under the control of the simian virus 40 promoter (P\text{SV40}). The L-arginine-responsive promoter (P\text{ART1}) harbors an ArgR-specific operator sequence (O\text{ARG}, capital letters, ARG boxes underlined) upstream of the minimal version of the cytomegalovirus immediate early promoter (P\text{hCMVmin}) and drives expression of a reporter gene (e.g. human secreted alkaline phosphatase, SEAP) in a L-arginine-induced manner. All expression units are terminated by a polyadenylation site (pA). Selected restriction sites are indicated: A, NcoI; B, BssHII; Ba, BamHI; E, EcoRI; H, HindIII; N, NotI; S, SfiI; X, XhoI, Xh, XhoI. (B) At a low L-arginine concentration (10 mg/l), the ARG1 is in a low-affinity DNA binding state and does not interact with its specific operator sequence (O\text{ARG}); therefore, expression of the transgene remains silent. At a higher L-arginine concentration (1 g/l), the ARG1 is in a high-affinity DNA binding state and interacts with its specific operator sequence (O\text{ARG}) and activates transcription from P\text{ART1}.

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CHO-K1 cells co-transfected with pSH91 (pSH91, PSV40-ARG2-pA). In general, increasing the number of operator modules resulted in higher maximum expression levels but also higher basal expression (Figure 4A). Having defined the parameters for optimal transgene regulation, the ART system was further validated for expression of the multiple sclerosis therapeutic interferon-β. Interferon-β was cloned downstream of P_{ART1} (pSH113, P_{ART1}-INF-β-pA), which was transactivated by ARG2 (pSH91, PSV40-ARG2-pA). CHO-K1 transiently co-transfected with pSH113 and pSH91 enabled adjustable INF-β expression, when exposed to l-arginine concentrations ranging from 10 mg/l to 1 g/l (Figure 4B).

Validation of the ARG2—P_{ART1} configuration in different cell lines

Since the combination of ART2 and P_{ART1} provided the best combination of low leaky and maximum expression levels, we sought to further validate this system. Immortalized cell lines of human, rodent and monkey origin were co-transfected with pSH91 and pSH93. Sixty hours post transfection, cells exposed to 10 mg/l L-arginine displayed low SEAP expression, whereas 1 g/l L-arginine induced high-level production of the reporter gene (Table 2). Moreover, expression integrity of ARG2 in mammalian cells was confirmed by western blot analysis using HEK293-T transiently transfected with pSH91 (PSV40-ARG2-pA) (Figure 5).

Autoregulated l-arginine-inducible transgene expression in CHO-K1

The ART system was also validated in an autoregulated positive feedback configuration enabling one-step installation of regulated transgene expression in mammalian cells using a single-vector format. This configuration mediates simultaneous expression of transactivator and transgene, both driven by the transactivator-dependent promoter and was found to be instrumental for the design of noise resistant gene networks (58). Leaky transscripts mediated by the P_{ART1} promoter initiate production of relatively few transactivator molecules which trigger full expression of P_{ART1}-driven transgenes in the presence of l-arginine. SEAP expression was assessed in CHO-K1 transiently transfected with pSH122 (pSH122, P_{ART1}-SEAP-IRE-PSV-ARG2-pA, Figure 6A). P_{ART1} transcription mediates basal SEAP and ARG2 production which results in ARG2-triggered auto-induction of the dicstrionic expression unit in the presence of high l-arginine concentrations (Figure 6B).

(PSV40). Selected restriction sites are indicated: B, BstHII; Ba, BamHI; E, EcoRI; H, HindIII; N, NotI; Sa, SalI; X, XbaI. The l-arginine-responsive promoters harboring 0 (P_{ART1}), 2 (P_{ART2}), 4 (P_{ART4}), 6 (P_{ART6}), 8 (P_{ART8}) and 10 (P_{ART10}) bp linkers between the operator sequence and the minimal promoter were co-transfected with l-arginine-dependent transactivators containing the (B) human NF-κB (ARG1, ArgR-κB), (C) human E2F4 (ARG3, ArgR-E2F4) or H. simplex VP16 (ARG2, ArgR-VP16). The expression of both transactivators is driven by the simian virus 40 promoter.

**Figure 3.** Regulation performance of P_{ART1} variants with different l-arginine-dependent transactivators. (A) Schematic representation of the different l-arginine-dependent transactivators. The bacterial repressor ArgR of Chlamydia pneumoniae, fused to human E2F4 (ARG3, ArgR-E2F4) or H. simplex VP16 (ARG2, ArgR-VP16). The expression of both transactivators is driven by the simian virus 40 promoter.
Table 2. Quantitative SEAP expression profiles under the control of \( \text{P}_{\text{ART1}} \) in immortalized cell lines with pSH91, an ARG2 encoding vector and pSH93, a \( \text{P}_{\text{ART1}} \)-driven SEAP expression cassette, SEAP expression levels were determined in cell culture media (U/l), 60 h after co-transfection

| Cell line  | \( 10 \text{mg/l L-arginine} \) | \( 1 \text{g/l L-arginine} \) |
|-----------|-------------------------------|-------------------------------|
| CHO-K1    | 0.69 ± 0.45                   | 8.08 ± 0.80                   |
| HEK-293T  | 1.67 ± 0.25                   | 21.83 ± 3.19                  |
| NIH/3T3   | \((0.72 ± 0.39) \times 10^{-3}\) | \((25.05 ± 2.00) \times 10^{-3}\) |
| COS-7     | 0.15 ± 0.07                   | 2.47 ± 0.20                   |
| HT-1080   | 0.04 ± 0.00                   | 0.72 ± 0.06                   |

Figure 4. (A) Validation of \( \text{P}_{\text{ART1}} \) variants containing a different number of ARG-specific operator modules. SEAP expression vector encoding L-arginine-responsive promoters harboring monomeric (pSH117, \( \text{P}_{\text{ARTm1}-\text{SEAP-pA}} \); \( \text{P}_{\text{ARTm1}} \), \text{Asc}I-OARG-MluI-0bp-\text{PhCMVmin})), dimeric (pSH119, \( \text{P}_{\text{ARTm2}-\text{SEAP-pA}} \); \( \text{P}_{\text{ARTm2}}, \text{Asc}I-OARG-7bp-OARG-MluI-0bp-\text{PhCMVmin})), trimeric (pSH126, \( \text{P}_{\text{ARTm3}-\text{SEAP-pA}} \); \( \text{P}_{\text{ARTm3}}, \text{Asc}I-OARG-7bp-OARG-7bp-OARG-MluI-0bp-\text{PhCMVmin})), or tetrameric (pSH127, \( \text{P}_{\text{ARTm4}-\text{SEAP-pA}} \); \( \text{P}_{\text{ARTm4}, \text{Asc}I-OARG-7bp-OARG-7bp-OARG-7bp-OARG-MluI-0bp-\text{PhCMVmin}}) \) operator modules were co-transfected with pSH91 (P\text{SV40-ARG2-pA}) into CHO-K1 and SEAP production was profiled after 60 h. The induction factor is shown on the top of each bar. (B) Dose–response profile of interferon-\( \beta \) expression in CHO-K1. Cells were transiently co-transfected with pSH113 (pSH113, \( \text{P}_{\text{ART1-INF-\beta-pA}} \)) and pSH91 (P\text{SV40-ARG2-pA}) and grown for 48 h at different L-arginine concentrations before quantification of the interferon-\( \beta \) production in the supernatant. Fold induction is shown on the top of each bar.

Figure 5. Western blot analysis of ARG2 expression in HEK-293T cells transfected with pSH91 (pSH91, P\text{SV40-ARG2-pA}) and cultivated for 60 h. Lane 1, lysate from HEK-293T transfected with pSH91 (P\text{SV40-ARG2-pA}); lane 2, lysate from untransfected control cells. The 35 kDa band, indicative for the fusion protein ARG2 is shown with a black arrow. The loading control (tubulin-a, 57kDa) is indicated with an open arrow. Migration of molecular mass markers (kDa) is indicated on the left.

Figure 6. (A) Diagram of the autoregulated L-arginine-inducible SEAP expression vector (pSH122). pSH122 harbors the L-arginine-responsive promoter (\( \text{P}_{\text{ART1}} \)) which drives transcription of the dicistronic expression unit encoding the the human placental alkaline phosphatase (SEAP) in the first and the L-arginine-dependent transactivator (ARG2) in the second cistron. Whereas translation of SEAP occurs via a classic cap-dependent mechanism, translation-initiation of ARG2 is mediated by an internal ribosome entry site of polioviral origin (IRESV). pA is the polyadenylation signal. Selected restriction sites are indicated: A, AatII; E, EcoRI; N, NotI; X, XbaI, Xh, XhoI. (B) pSH122 was transiently transfected into CHO-K1, cultivated in media containing 10 mg/l or 1 g/l L-arginine prior to SEAP quantification. Fold induction is indicated on the top of the bars.
Regulation performance of L-arginine derivatives and secondary products

In order to assess the specificity of ART-controlled transgene expression, we cultivated CHO-K1, co-transfected with pSH91 (Psv40-ARG2-pA) and pSH93 (PART1-SEAP-pA), in the presence of increasing concentrations of L-arginine and several of its derivatives (L-canavanine, L-homoarginine, L-arginine methyl ester and L-arginine ethyl ester) and secondary metabolic products (L-ornithine, L-citrulline, agmatine) and profiled SEAP production after 60 h (Figure 7A). Of all these compounds only L-arginine and its methyl and ethyl esters induced SEAP to significant levels while retaining typical dose–response characteristics (Figure 7B).

Adjustability and reversibility of the ART-controlled transgene expression

In order to determine the adjustability and reversibility of L-arginine-controlled transgene expression, we generated a stable CHO-K1-derived cell line transgenic for constitutive ARG2 expression and P ART1-controlled SEAP production (CHO-ARG2-SEAP). CHO-ARG2-SEAP was cultivated for 60 h in medium containing increasing concentrations of L-arginine before SEAP was profiled in the culture supernatant. As shown in Figure 8A, SEAP expression could be precisely adjusted to specific levels which correlate with particular L-arginine doses. Besides adjustability, rapid response kinetics and reversibility are key assets for a mammalian transgene control system. In order to assess the expression kinetics of the ART system, we profiled both SEAP expression and cell growth for over 96 h. CHO-ARG2-SEAP cells cultivated in medium supplemented with either 10 mg/l or 1 g/l L-arginine displayed similar growth rates, whereby significant SEAP expression was detected only in CHO-ARG2-SEAP cultivated in the presence of 1 g/l L-arginine (Figure 8B). The reversibility was determined by cultivating CHO-ARG2-SEAP for up to 1 week while alternating the inducing (1 g/l) or repressing (10 mg/l) culture condition every other day. The SEAP production profiles revealed reproducible sequential expression kinetics, precise reversibility and tight repression after ON-to-OFF switching (Figure 8C).

L-arginine-inducible transgene expression in mice

For validation of ART-controlled transgene expression in vivo, we have encapsulated CHO-ARG2-SEAP into coherent alginate-poly-(L-lysine)-alginate microcapsules (200 cells per capsule) and implanted them (10^6 capsules per mouse) intraperitoneally into mice. Implanted mice were given a daily dose of L-arginine ranging from 0 to 100 mg/kg for 72 h before quantifying serum SEAP levels. The SEAP levels reached in the serum of treated mice was adjustable and dependent on the injected L-arginine dose. The serum SEAP levels of control mice implanted with microencapsulated CHO-K1 (data not shown) or mice implanted with microencapsulated CHO-ARG2-SEAP but not receiving exogenous L-arginine showed insignificant SEAP expression level (Figure 9A). In parallel, the experiment was also performed in vitro with the same batch of encapsulated cells, which confirmed the dose–response characteristics of microencapsulated CHO-ARG2-SEAP (Figure 9B).

DISCUSSION

Currently available transgene control systems have been designed for optimal regulation performance in vitro or in...
To enable interference-reduced operation within a complex biochemical reaction network of the host cell, heterologous or modified endogenous transcription factors have been engineered to modulate transcription of specific target promoters (60,61). Initially, inducer molecules only required reasonable pharmacokinetics and needed to be non-toxic at regulation-effective concentrations, criteria which was best met by clinically licensed small-molecule drugs (16,18,62). However, the use of clinically licensed substances to control biopharmaceutical manufacturing of difficult-to-produce protein pharmaceuticals fails to comply with administrative regulations or requires prohibitive downstream processing standards (33,63,64). Also, long-term administration of drugs at (sub-)clinical doses has been associated in many cases with side-effects which limits their use in future gene therapy trials (26,28–30). In contrast, physiologic molecules such as amino acids are FDA-licensed media components and an integral part of host metabolism. Due to possible interference effects, such physiologic trigger molecules have so far not been considered for transgene control.

As an obligate intracellular pathogen, Chlamydia pneumoniae has evolved to optimally plug in its metabolism into the biochemical networks of host cells (38).
As a major part of this metabolic crosstalk, the biosensor ArgR quantifies L-arginine levels and manages import of this amino acid (40). It is, therefore, reasonable to assume that the sensitivity of ArgR is attuned to physiologic L-arginine levels of mammalian cells. Indeed, ART-controlled expression of reporter and product genes shows excellent regulation performance (rapid induction kinetics, full adjustability, reproducible reversibility) in different cell lines as well as in mice with an arginine concentration window between 10 mg/l (required to maintain metabolism, transgene repressed) and 1 g/l (full induction). The L-arginine level in the plasma of mice is around 17 mg/l, which is well below the threshold concentration required to fully activate the ART system (52). By default, ART is an ON-type system, which means that it is only induced in the presence of increased L-arginine levels and remains repressed at physiologic concentrations of this amino acid.

ART is exclusively regulated by L-arginine and its ester forms which are rapidly converted to L-arginine by endogenous esterases (65). Secondary products of the L-arginine metabolism or the synthetic L-homoarginine failed to regulate the ART system. Even L-canavanine, which is stereochemically similar to L-arginine and was shown to fit in the amino acid binding pocket of ArgR was unable to modulate ART-controlled transgene expression (66).

Since L-arginine is a licensed component of cell culture media, the ART system will be a extremely valuable tool for the biomanufacturing of therapeutic proteins. Moreover, L-arginine is considered to be compatible with prolonged therapeutic use. In fact, L-arginine is currently therapeutically administered for pulmonary hypertension of newborns (67). It also improves endothelial function in animal models of hypercholesterolemia and atherosclerosis with no life-threatening side-effects reported so far (68).

As a prototype transgene control system responsive to a physiologic trigger molecule, the ART system has shown that metabolic and synthetic control circuits can functionally co-exist while sharing the same inducer. Although such systems may be of immediate use in gene-function analysis, gene network design as well as for biopharmaceutical manufacturing, the ART may develop its highest impact in future therapies: synchronization of therapeutic gene expression with fluctuations of endogenous molecules or the self-sufficient treatment of a pathological situation by exploiting signal molecules for a self-controlling genetic intervention are highly appealing strategies for future gene therapy applications.

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