Microtubule-associated protein tau facilitates the targeted killing of proliferating cancer cells \textit{in vitro} and in a xenograft mouse tumour model \textit{in vivo}

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\textbf{Background:} Antibody drug conjugates (ADCs) and immunotoxins (ITs) are promising anticancer immunotherapeutics. Despite their encouraging performance in clinical trials, both ADCs and ITs often suffer from disadvantages such as stoichiometrically undefined chemical linkage of the cytotoxic payload (ADCs) and the potential immunogenicity of toxins derived from bacteria and plants (ITs).

\textbf{Methods:} Human microtubule-associated protein tau (MAP) was cloned in-frame with human EGF, expressed in \textit{E. coli} and purified by standard chromatographic methods. The \textit{in vitro} activity was confirmed by flow cytometry, cell viability assays and tubulin polymerisation assay. The \textit{in vivo} efficacy was demonstrated using noninvasive far-red \textit{in vivo} imaging.

\textbf{Results:} The EGF-MAP selectively induced apoptosis in EGFR-overexpressing proliferating cancer cells through stabilisation of microtubules. Nonproliferating cells were not affected, demonstrating superior selectivity of EGF-MAP for cancer cells. The EGF-MAP was well tolerated at high doses in mice compared with the ETA'-based control. The \textit{in vivo} efficacy of EGF-MAP was demonstrated in a tumour xenograft mouse model.

\textbf{Conclusion:} Our data indicate the general mechanism of action for a new class of human immunotherapeutic reagents suitable for the treatment of cancer. This approach combines the binding specificity of targeting ligands with the selective cytotoxicity of MAP towards proliferating cells.

One prominent goal in passive immunotherapy is the targeted delivery of highly cytostatic payloads to malignant cells. This can be achieved using bifunctional molecules, comprising a targeting component and a cytotoxic component. The targeting component can be a full-size monoclonal antibody (mAb), a fragment thereof or a protein ligand (e.g., a growth factor or cytokine) (Cawley \textit{et al}, 1980; Williams \textit{et al}, 1987; von Minckwitz \textit{et al}, 2005). A variety of substances have been used as cytotoxic agents, including radioisotopes, small-molecule drugs and protein-based toxins derived from bacteria or plants. Depending on the linkage between the targeting and cytotoxic components, these molecules are described as antibody drug conjugates (ADCs) with chemical crosslinking between synthetic cytotoxic molecules and antibodies, or immunotoxins (ITs) if cytotoxic proteins are genetically fused to the targeting protein (Wu and Senter, 2005; Pastan \textit{et al}, 2006; Carter and Senter, 2008). Both ADCs and ITs can overcome the
disadvantages of naked mAbs, which often require a functional host immune system for their therapeutic effect (Weiner et al., 2010).

At least 22 ADCs are currently undergoing clinical trials. Two ADCs, Kadcyla (ado-trastuzumab emtansine) and brentuximab vedotin, have already been approved by the FDA for the treatment of HER2- and CD30-positive cancers, respectively (Lambert, 2005; Vedotin, 2012; Sievers and Senter, 2013). Interestingly, the cytotoxic domain of brentuximab vedotin is monomethyl auristatin E, a synthetic analogue of the tubulin polymerisation inhibitor dolastatin 10 (Tanaka et al., 2009). The success of this ADC underlines the enormous potential of cytostatic agents delivered by targeted immunotherapy. Classically, the cytotoxic compound is conjugated to the targeting mAb via nondirected chemical linkage, often resulting in an undefined stoichiometry of ADCs and heterogenic positioning of the cytotoxic payload in relation to the binding protein, which can directly affect its therapeutic efficacy (Ducry and Stump, 2010). Similarly, one disadvantage of chimeric ITs is their combination of a human or humanised binding component (e.g., a single-chain antibody fragment, scFv) and a toxin derived from bacteria or plants that can induce undesirable immune responses and dose limitations, for example, because of vascular leak syndrome (Li et al., 2009). The cytotoxic domain of brentuximab vedotin is monomethyl auristatin E, a synthetic analogue of the tubulin polymerisation inhibitor dolastatin 10 (Tanaka et al., 2009). The success of this ADC underlines the enormous potential of cytostatic agents delivered by targeted immunotherapy.

Human RNases 1, 2, 3 and 5 (angiogenin), apoptotic human enzymes in the context of fully human cytolytic fusion proteins (CFPs). Human RNases 1, 2, 3 and 5 (angiogenin), which degrade RNA and induce apoptosis by inhibiting protein synthesis, have been used to replace non-human toxins (Mathew and Verma, 2009). The specific cytotoxicity of human angiogenin towards CD30-overexpressing Hodgkin’s lymphoma-derived cell lines has been demonstrated by fusion with the CD30 ligand (Huhn et al., 2001). Another CD30-targeted CFP was shown to kill CD30-overexpressing tumour cells efficiently when combined with human death-associated protein kinase 2 (Tur et al., 2009). In 2008, a completely human granzyme B-based CFP directed against CD64 was found to be toxic towards an acute myeloid leukemia (AML)-related cell line and primary AML cells (Stahnke et al., 2008). Thus far, various human proteins including RNases, perforin, Bik, Bak, Bax, DNA fragmentation factor 40, FAS ligand and TNF-related apoptosis-inducing ligand have been used as toxic effector molecules (ten Cate et al., 2010). Although most of these have been tested for toxicity towards cancer cells, they could potentially be applied in other indication areas such as autoimmune and chronic inflammatory disorders.

Generally, only the binding domain of CFPs confers selectivity and specificity, whereas the cytotoxic component is chosen according to its ability to induce apoptosis based on different modes of action. However, most tumour markers are also expressed on physiologically normal cells, resulting in undesirable off-target effects. Cytostatic agents such as taxanes, which degrade RNA and induce apoptosis by inhibiting protein synthesis, have been used to replace non-human toxins (Mathew and Verma, 2009). The specific cytotoxicity of human angiogenin towards CD30-overexpressing Hodgkin’s lymphoma-derived cell lines has been demonstrated by fusion with the CD30 ligand (Huhn et al., 2001). Another CD30-targeted CFP was shown to kill CD30-overexpressing tumour cells efficiently when combined with human death-associated protein kinase 2 (Tur et al., 2009). In 2008, a completely human granzyme B-based CFP directed against CD64 was found to be toxic towards an acute myeloid leukemia (AML)-related cell line and primary AML cells (Stahnke et al., 2008). Thus far, various human proteins including RNases, perforin, Bik, Bak, Bax, DNA fragmentation factor 40, FAS ligand and TNF-related apoptosis-inducing ligand have been used as toxic effector molecules (ten Cate et al., 2010). Although most of these have been tested for toxicity towards cancer cells, they could potentially be applied in other indication areas such as autoimmune and chronic inflammatory disorders.

Cloning of EGF-MAP. The open reading frame (ORF) for human EGF (GeneID: 1950) was modified with SfiI and NotI restriction sites by PCR followed by ligation into a SfiI/NotI-linearised pMT vector. The ORF for MAP (Gene ID: 4137; NM_016834.4) was modified with SfiI and NotI-linearised vector already containing the EGF sequence (Figure 1A). Two point mutations were introduced to remove phosphorylation sites by PCR followed by ligation into a NotI/BglII-linearised pMT vector compatible solutes as described previously (Barth et al., 2000). Briefly, bacteria were grown after transformation to an OD of 1.6 followed by stress induction with 500 mM D-sorbitol, 10 mM betaine monohydrate and 4% (w/v) NaCl. After incubation for 30 min at 26 °C with shaking (180 r.p.m.), protein expression was induced with 2 mM IPTG. Bacteria were harvested 18 h after induction by centrifugation (4000 × g, 10 min, 4 °C) and frozen at –80 °C overnight. The frozen pellet was resuspended in preparation buffer (75 mM Tris-HCl, 300 mM NaCl, 5 mM DTT, 10 mM EDTA, 10% (v/v) glycerol, pH 8.0) containing a complete protease inhibitor cocktail (Roche, Mannheim, Germany) at 4 °C and

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sonicated 5 times for 60 s at 200 W. Cell debris was removed by centrifugation (24 000 × g, 20 min, 4 °C) and EDTA was removed by dialysis against PBS (pH 7.4) at 4 °C overnight. The EGF-MAP was purified by IMAC on an Äkta Purifier System (GE Healthcare, by dialysis against PBS (pH 7.4) at 4 °C/mg and SEC. Preparative SEC was carried out using an Äkta Purifier System (GE Healthcare) with a XK16–70 column packed with Superdex 75 (GE Healthcare), and a mobile phase of PBS (pH 7.4) at a flow rate of 1.0 ml min⁻¹.

**SDS–PAGE and western blot analysis.** Protein enrichment and identity were determined by sodium dodecylsulphate polyacrylamide gel electrophoresis (SDS–PAGE) followed by Coomassie brilliant blue staining and western blotting, respectively, as previously described (Hetzel et al, 2008). The protein concentration was determined by densitometry against bovine serum albumin standards using AIDA software (Raytest GmbH, Straubenhardt, Germany) after staining with Coomassie brilliant blue, and was verified using the Bradford assay (Bio-Rad, Munich, Germany). The EGF-MAP was detected by western blot using mouse-anti-penta-His (1 : 5000, Sigma, Germany) and western blot analysis of EGF-MAP. The EGF-MAP was purified by IMAC and SEC. The enrichment was determined by SDS–PAGE followed by staining with Coomassie brilliant blue (lane 1). Protein identity was verified by western blotting (lane 2) using mouse-anti-penta-His (1 : 5000, Qiagen). The primary antibody was detected using an alkaline phosphatase-conjugated anti-mouse-IgG mAb (1 : 5000; Sigma, Germany) followed by staining with NBT/BCIP substrate (Life Technologies). (C) The EGF-MAP cell-binding analysis. Purified EGF-MAP was tested for its ability to bind EGFR + L3.6pl cells (EGFR–HEK293 cells were used as a negative control). The 425(scFv)-ETA' has previously been shown to bind L3.6pl cells and was used as a positive control. Bound EGF-MAP and 425(scFv)-ETA' were detected with anti-penta-His-Alexa Fluor 488 antibody (Qiagen).

**Transfection of L3.6pl cells with Katushka-2.** Transfection was carried out as described previously (Pardo et al, 2012). Briefly, we seeded 1 × 10⁵ L3.6pl cells in a 12-well plate and transfected them with DNA encoding the far-red fluorescent protein Katushka-2 (pTag-Katushka2-N; Evrogen, Heidelberg, Germany) using Roti-Fect (Roti, Karlsruhe, Germany) according to the manufacturer’s instructions. Transfected cells were selected with G418 (InvivoGen, Toulouse, France) and sorted using the FACS Vantage Cell Sorter (Becton Dickinson, Heidelberg, Germany).

**Binding analysis by flow cytometry.** The cell-binding activity of purified EGF-MAP was analysed by flow cytometry. We incubated 4 × 10⁵ cells with 1 μg EGF-MAP in PBS (pH 7.4) containing 2 mM EDTA and 0.5% (w/v) BSA for 30 min on ice followed by washing with PBS. Fluorescence staining was performed using an anti-penta-His-Alexa Fluor 488 antibody (1 : 100; Qiagen, Germany) for 30 min on ice in the dark. Finally, the cells were washed twice with PBS and analysed on a FACS Calibur flow cytometer (Becton Dickinson, Heidelberg, Germany).

**In vitro cellular cytotoxicity.** The cytotoxic effect of EGF-MAP was assessed by measuring the conversion of XTT to a water-soluble orange formazan dye. We seeded 1 × 10⁶ cells per well into a 96-well microtitre plate and incubated the cells with various dilutions of the recombinant protein for 72 h at 37 °C, 5% CO₂ and
100% humidity. We used 425(scFv)-ETA', human recombinant EGF and the EGFR⁺ cell line HEK293 controls. We added 50 μl XTT/phenazine methosulphate (100:1; Serva and Sigma, Steinheim, Germany) to each well and incubated the plates as above for 3–4 h before measuring the absorbance at 450 and 630 nm using an Epoch Microplate Spectrophotometer (Biotek, Bad Friedrichshall, Germany). The concentration required to achieve 50% reduction of protein synthesis (IC₅₀) relative to untreated control cells was calculated using GraphPad Prism 5 (GraphPad Software, La Jolla, CA, USA). All experiments were carried out in triplicate.

**Apoptosis assay.** An AnnexinV/propidium iodide assay was used to determine the pro-apoptotic impact of EGF-MAP. We incubated 2.5 × 10⁶ cells per ml with 100 nM EGF-MAP in a 12-well plate (Greiner, Frickenhausen, Germany) for 48 h at 37°C, 5% CO₂ and 100% humidity. We used 425(scFv)-ETA' (10 nM) and EGFR⁺ HEK293 cells as controls. After incubation, the cells were washed twice with PBS (pH 7.4) and stained with AnnexinV-FITC (ebioscience, Frankfurt, Germany) in AnnexinV binding buffer (10 mM HEPES/NaOH (pH 7.4), 140 mM NaCl, 2.5 mM CaCl₂) for 30 min at room temperature. Finally, the cells were washed again as described above, resuspended in AnnexinV buffer containing 10 μg ml⁻¹ propidium iodide and analysed by flow cytometry using a FACS Calibur (Becton Dickinson, Germany).

**Tubulin polymerisation assay.** The stabilising effect of EGF-MAP on tubulin polymerisation was tested using a Tubulin Polymerisation Assay Kit (tebu-bio, Offenbach, Germany) according to the manufacturer’s instructions. Paclitaxel and general tubulin tubulin were used as controls. All measurements were carried out in duplicate. Vmax represents the slope of the linear phase in milli-extinction (mE) per min.

**In vivo analysis.** Animal experiments were officially approved by the local Animal Care and Use Review Committee. All animals received humane care in accordance with the requirements of the German Tierschutzgesetz, §§ Abs. 1 and in accordance with the guide for the care and use of laboratory animals published by the National Institutes of Health in 2011. Animal experiments were carried out as described previously (Pardo et al, 2012). Briefly, 1 × 10⁶ L3.6pl cells expressing Katushka-2 were suspended in ~20 μl PBS (pH 7.4) and subcutaneously injected into the right hind limb of 6–8-week-old female Balb/C nu/nu mice (Charles River, Sulzfeld, Germany). For imaging experiments, mice were placed on a purified, chlorophyll-free diet (AIN93G, SSNIFF GmbH, Soest, Germany) 11 days before the imaging experiments began. The animals were injected intravenously with EGF-MAP (4 mg kg⁻¹) or PBS (pH 7.4) according to the treatment regimen shown in Figure 6. Readouts were taken on the same days as the treatment, using the Maestro CRI optical imaging system (Maestro CRI Inc., Waltham, MA, USA). Images were analysed using the Maestro spectral imaging software as previously described (Kampmeier et al, 2010). The Katushka-2 signal was detected using the yellow filter set (630–850 nm).

**Statistical analysis.** Statistical analysis was performed using the GraphPad Prism 5 (GraphPad Software). Data were expressed as the mean ± s.d. or s.e.m. as indicated. Statistical comparisons were made using a two-tailed unpaired Student’s t-test. *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001.

**RESULTS**

Generation of EGF-MAP and characterisation of binding activity. We considered a range of candidate effector molecules for CFPs based on the need to use a fully human sequence with a low molecular weight suitable for fusion proteins, and with the ability to induce apoptosis in target cells without nonspecific toxicity to healthy cells (e.g., by nonspecific surface binding). Human MAP met these criteria. The protein exists in eight isoforms, all derived from a single precursor RNA by alternative splicing. We selected isoform 3 because this was the smallest protein that retained all four highly conserved microtubule-binding repeats.

In contrast to the hyperphosphorylated tau protein that can form paired helical filaments and neuronal tangles associated with Alzheimer’s disease (Morris et al, 2011), MAP is unable to aggregate. However, the MAP amino acid sequence includes numerous potential phosphorylation sites, many of which regulate its activity. We removed two phosphorylation sites, S156 and S204, by site-directed mutagenesis, which have been shown to critically regulate microtubule binding (Schneider et al, 1999). Both serine residues were replaced with alanine by PCR. Furthermore, we added a C-terminal NLS derived from the SV40 T-antigen by adding the sequence (5'–cccccccccccccccccccccccgt-3') to the 3' end of the ORF to foster nuclear enrichment of the internalised protein. The complete ORF was then transferred to the pMT vector (Figure 1A) linearised with NotI/BbI, placing it in frame with an upstream ORF representing human EGF.

The construct was introduced into E. coli BL21 (DE3) cells and EGF-MAP was expressed at yields of up to 1 mg of purified protein per litre of bacterial culture. After one IMAC purification step and SEC, the enrichment and identity of EGF-MAP were determined by SDS–PAGE following by Coomassie brilliant blue staining and western blotting, respectively (Figure 1B). Although purity was low, subsequent functional analyses could be carried out always including appropriate negative controls. Specific binding activity to L3.6pl cells was confirmed by flow cytometry, which also revealed the absence of nonspecific binding to the EGFR⁺ HEK293 cells (Figure 1C).

EGF-MAP shows dose- and proliferation-dependent cytotoxicity towards L3.6pl cells. The **in vitro** cytotoxicity of EGF-MAP was determined using an XTT cell viability assay. The EGF-MAP showed specific toxicity towards L3.6pl cells with an IC₅₀ value of 1 μM, whereas the EGFR⁺ cell line HEK293 was unaffected (Figure 2). Recombinant human EGF was used as control with no adverse effects on cell viability. In addition to our model cell line L3.6pl, EGF-MAP was cytotoxic to other cancer cell lines expressing different levels of EGFR (Table 1). As we used EGF-MAP as a model protein to demonstrate the cytotoxic potential of MAP as an effector, we were not interested in establishing an elaborated purification protocol for this fusion protein, which might lead to underestimated IC₅₀ values.

To test our hypothesis that MAP acts in a proliferation-dependent manner, we generated another fusion protein comprising MAP in combination with the EGFR-specific antibody fragment 425(scFv). The 425(scFv) is an EGFR-blocking antibody fragment. Antibodies that block EGFR have previously been shown to induce G₁ arrest in EGFR-overexpressing cell lines, thus blocking proliferation (Kiyota et al, 2002). When we tested the 425(scFv)-MAP construct, we observed the binding to L3.6pl cells (data not shown) but no cytotoxicity (Figure 3A), confirming that the cytotoxicity of MAP is proliferation dependent. In contrast, the control immunotoxin 425(scFv)-ETA' was highly cytotoxic to L3.6pl cells (IC₅₀ = 1 μM) even in their G₁ arrested state (Figure 3A). To confirm this finding, we induced mitotic block in L3.6pl cells using a previously pre-titrated cytostatic but non-apoptosis-inducing (Pasquier et al, 2004) concentration of paclitaxel (100 pM for 24 h) followed by the addition of 1 μM EGF-MAP. There was no cytotoxicity of EGF-MAP to nonproliferating L3.6pl cells, whereas apoptosis was strongly induced in L3.6pl cells that have not received paclitaxel before (Figure 3B).
EGF-MAP cytotoxicity involves the induction of apoptosis. The EGF-MAP mechanism of action was investigated by incubating L3.6pl cells with EGF-MAP followed by staining with Annexin V-FITC and propidium iodide. As this assay was performed to demonstrate qualitatively the pro-apoptotic effect of the EGF-MAP, the concentration (100 nM for EGF-MAP and 10 nM for 425(scFv)-ETA’) and the incubation time (72 h) were appropriately chosen to capture the cells within the early and late apoptotic stage. The population of early and late apoptotic cells (lower and upper right corner) increased significantly, demonstrating that the cytotoxicity of EGF-MAP towards L3.6pl cells reflects the induction of apoptosis (Figure 4A). There was comparatively little induction of apoptosis in HEK293 cells (Figure 4B).

EGF-MAP stabilises microtubules in targeted cells. Apoptosis can be induced by modifying the polymerisation dynamics of microtubules, either by stabilising (e.g., paclitaxel) or destabilising them (e.g., nocodazole). The effector protein MAP was anticipated to interfere with mitosis by binding to and stabilising microtubules. To determine the mechanism of action, we carried out a tubulin polymerisation assay. A total of 1 μM (≈IC_{50}) EGF-MAP was able to stabilise microtubules effectively and promoted polymerisation, as reflected by the increased polymerisation velocity Vmax. As a control, 10 μM paclitaxel (concentration recommended by the manufacturer) was used (Figure 5).

EGF-MAP is well tolerated and inhibits tumour growth in vivo. Before treatment experiments, EGF-MAP was tested in a dose-escalation study in tumour-free mice to determine the maximum tolerated dose. Three mice per group received 0.5, 1.0, 2.0 or 4.0 mg kg^{-1} of EGF-MAP intravenously on days 0, 1, 3 and 5. The overall health status and body weight of the mice was monitored daily. All mice survived and showed normal behaviour. No reduction of body weight was observed (Figure 6A).

To test the efficacy of EGF-MAP in vivo, we transfected L3.6pl cells with the far-red fluorescent protein Katushka-2, and injected 1 × 10^6 transfected cells subcutaneously into the right hind leg of 10 Balb/C nu/nu mice to induce tumours. We monitored tumour growth using the in vivo imaging system Maestro Cri, based on the premise that the total Katushka-2 signal correlates with the cell number. We tested the injected mice 11 days after inoculation with tumour cells, and all mice showed palpable tumours (Figure 6B). Mice were then randomly divided into two groups (n = 5) that were allocated one treatment cycle with EGF-MAP (4 mg kg^{-1}) or PBS (pH 7.4). Data are presented as means ± s.e.m. Statistical analysis was carried out using a two-tailed unequal Student’s t-test (*P≤0.05, **P≤0.01, ***P≤0.001). Our results clearly demonstrated that EGF-MAP caused a significant reduction in tumour growth compared with the PBS control group (Figure 6C and D).

| Cell line | EGF-MAP | 425(scFv)-ETA’ | EGFR level |
|-----------|---------|----------------|------------|
| L3.6pl    | 1.0     | 0.001          |            |
| PC-3      | 2.5     | 0.002          |            |
| C4-2      | 2.8     | 0.04           |            |

Abbreviations: EGF = epidermal growth factor; EGFR = epidermal growth factor receptor; IC_{50} = half-maximal inhibitory concentration; MAP = microtubule-associated protein tau; 425(scFv)-ETA’ = single-chain fragment (425(scFv) genetically fused to a truncated version of Pseudomonas Exotoxin A.

**DISCUSSION**

Progress towards the development of therapeutically effective ADCs, ITs and CFPs depends on the selection of an appropriate target and the corresponding binding component, on the efficacy of the cytotoxic payload and on the nature of the linkage between these two modalities. Most ADCs currently undergoing clinical evaluation contain small synthetic cytotoxic molecules, such as maytansinoids or auristatins (Sievers and Senter, 2013). The architecture of ITs is exclusively chimeric, comprising a protein ligand chemically or genetically linked to a cytotoxic protein (Kreitman et al, 2001, 2009). Both formats often suffer from inherent limitations that can either be encountered during synthesis (in the case of ADCs, where non-directed coupling
produces heterogeneous products) or during testing (in the case of ITs, where non-human proteins such as ETA induce undesirable immune responses). To address these challenges, we have developed a new fully human CFP containing human MAP as the cytotoxic effector that kills cancer cells in a proliferation-dependent manner. The MAP was selected as a potential cytostatic effector against rapidly proliferating cancer cells based on its ability to regulate mitosis. It binds to microtubules directly through a common structural motif of 18 amino acids called microtubule-binding repeat (MBRs). These MBRs have a positive net charge that facilitates the interaction with negatively charged residues in tubulin (Jho et al., 2010). Binding to spindle microtubules during mitosis finally promotes microtubule assembly and stability (Weingarten et al., 1975). The intrinsic proliferative nature of cancer cells makes them a suitable target for this new effector protein. However, in addition to the selectivity conferred by using a cytotoxic effector that only affects proliferating cells, the inclusion of a targeting component ensures that the fusion protein is delivered only to cells expressing the target antigen. We tested EGF-MAP using EGFR-overexpressing L3.6pl cells as a model, as previously used to evaluate EGFR-targeted ITs such as 425(scFv)-ETA’ (Bruell et al., 2005; Pardo et al., 2012). These in vitro tests confirmed the efficacious induction of apoptosis in L3.6pl cells. We then used in vivo optical imaging to determine the efficacy of EGF-MAP on the prevention of tumour growth in vivo (Kampmeier et al., 2010; Pardo et al., 2012). This approach allowed us to monitor the therapeutic effect of EGF-MAP in a single animal, thus reducing the number of total animals required per experiment. Tumour growth was significantly inhibited by EGF-MAP compared with the untreated control group. Compared with the maximum tolerated dose of 425(scFv)-ETA’ (1.0 mg kg$^{-1}$) (Bruell et al., 2005), a dose of 4.0 mg kg$^{-1}$ of EGF-MAP could be administered without adverse effects and a lethal dose was not reached. Cancer immunotherapy using protein-based therapeutics such as EGF-MAP will require repetitive administration, and thus the relatively high tolerability of CFPs make them superior to classical ITs.

Because ‘tau’ is also the name of a protein associated with Alzheimer’s diseases, it is important to emphasise that the MAP tau protein described herein is not able to form the pathologically relevant paired helical filaments and our proposed therapeutic strategy does not increase the risk of neurodegenerative disorders (Dickson, 2004; Feinstein and Wilson, 2005; Iovino et al., 2010; Morris et al., 2011). Because of removal of two critical phosphorylation sites, our MAP tau is not subject to hyperphosphorylation. Beyond that, the targeted nature of MAP-based CFPs prevents them from accumulating in the extracellular space of the central and peripheral nervous system.
The cytotoxic efficacy of human effector proteins is not yet comparable to that of bacterial ETA. The higher IC50 values we observed for MAP in our in vitro experiments partially reflects its lack of enzymatic activity (ETA is an ADP-ribosyl transferase that acts catalytically against its targets, whereas MAP binds stoichiometrically to microtubules). However, the cytotoxic activity of CFPs such as MAP could be improved by the insertion of an adapter sequence that could facilitate transfer of the effector molecule from the endosome to the cytosol (Hetzel et al., 2008). For example, it has been reported that only 3–4% of internalised ETA is ultimately translocated (Ogata et al., 1990). Improving the translocation efficiency could therefore significantly enhance the therapeutic potential of CFPs and make them even more suitable for clinical applications. Furthermore, we used the natural EGF ligand as the targeting component to provide a model suitable for testing of the new effector protein MAP. Potential differences in binding affinity between EGF and the anti-EGFR 425(scFv) might also contribute to disparities in efficacy. In regard to future development and improvements of EGFR-targeting CFPs, non-blocking anti-EGFR mAbs instead of EGF could also be used to improve the therapeutic efficacy of MAP.

Because of their human nature and the anticipated low immunogenicity, MAP-based CFPs could also be applied in indication areas beyond cancer. Non-life-threatening diseases, including many inflammatory and autoimmune disorders, still lack effective and targeted therapeutic approaches. The modular architecture of CFPs allows the rapid replacement of the binding component with new ligands, antibodies or antibody fragments, potentially allowing the targeting of proliferating inflammatory cells, such as macrophages and T cells (Hristodorov et al., 2012).

In conclusion, we have developed a new fully human CFP comprising the cytostatic effector MAP, allowing the induction of...
Apoptosis specifically in proliferating target cells. Using EGF ligand as the binding component and L3.6pl cells as a model, we have demonstrated the cytotoxic potential of EGF-MAP in vitro and in vivo. Based on our findings, MAP-based immunotherapeutic reagents appear to be promising candidates for targeted immunotherapy in cancer and other immunological diseases.

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**CONFLICT OF INTEREST**

DH, RM, TT and SB are co-inventors on a corresponding patent assigned to Fraunhofer. The other authors declare no conflict of interest.

**AUTHOR CONTRIBUTIONS**

Conception and design: D Hristodorov, A Pardo, M Huhn and S Barth; acquisition of data: D Hristodorov, A-T Pham and R Mladenov; analysis and interpretation of data: D Hristodorov; writing, review and/or revision of the manuscript: D Hristodorov, R Fischer, S Barth and T Thepen; animal study supervision: A Pardo and T Thepen.
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