Chemical modification allows phallotoxins and amatoxins to be used as tools in cell biology

Jan Anderl¹, Hartmut Echner² and Heinz Faulstich*³

Abstract
Phallotoxins inhibit the dynamics of microfilaments in cells and lead to apoptosis. Due to poor cellular uptake these effects cannot be studied in live cells, even at millimolar toxin concentrations, nor can phalloidin be used for the elimination of tumor cells. Uptake is greatly enhanced by conjugation of phallotoxins to either lipophilic or polycationic moieties, such as oleic acid, polylysine, or Tat-peptide. These conjugates were lethally toxic for cells, e.g., mouse fibroblasts or Jurkat leukemia cells, in the micromolar range. Uptake into cells starts with the attachment of the toxin conjugates to the plasma membrane, followed by endocytosis and, in most cases, cleavage of the toxin from the carrier. Interestingly, the internalization rate of phalloidin into cells was also significantly increased by the fluorescent moiety tetramethylrhodaminyl, as well as by high molecular weight methoxy-polyethylene-glycol, two compounds unknown so far for their uptake-mediating activity. Conjugation to carriers as investigated in this work will allow the use of phallotoxins in experimental cell biology and possibly in tumor therapy. The findings obtained with phallotoxins could be applied also to the family of amatoxins, where α-amanitin, for example, when conjugated to oleic acid was more than 100-fold more toxic for cells than the native toxin. This suggests the possibility of a more general use of the moieties examined here to enhance the uptake of hydrophilic peptides, or drugs, into live cells.

Introduction
Phallotoxins and amatoxins, the two families of toxic cyclopeptides produced by the green death cap Amanita phalloides, have been the subject of intense biochemical research for decades [1]. Although produced by the same mushroom and of similar structure, the two peptide families have totally different cellular targets. Phallotoxins, such as phalloidin, bind to polymeric actin, thus stabilizing microfilaments and decreasing the amount of monomeric actin in equilibrium with the filaments. This
interaction is in the nanomolar range and highly specific: no other targets for phalloidin in the cell are known. Amatoxins, such as the main toxin \( \alpha \)-amanitin, bind to RNA-polymerases II of eukaryotic cells, thus inhibiting the transcription process at nanomolar concentrations. Also this interaction is specific, since RNA-polymerases I are not inhibited at all, whereas RNA-polymerases III are inhibited at amanitin concentrations ca. 1000 times higher than for RNA-polymerases II [2,3]. The fact that both the cytoskeleton and the eukaryotic transcription machine are complex structures and still under investigation, may explain the continuing interest in these two kinds of specific inhibitors.

Phalloidin has been used to study actin dynamics in vitro [1], and in microscopic studies after microinjection into single cells [4]. Beside such experiments phalloidin conjugated to fluorescent moieties is widely used for visualizing filamentous actin in fixed cells [5,6]. In all these applications, cell-free systems were used, allowing direct access of phalloidin to its target. Similarly, \( \alpha \)-amanitin was mainly used with isolated nuclei or the solubilized enzyme, as recently reported for stabilizing yeast RNA-polymerase II in an X-ray analysis [7].

For both phallotoxins and amatoxins, experience with live cells is limited by the fact that the peptides cross the plasma membrane barrier only very slowly. Poor uptake rates of phallotoxins and amatoxins have been observed for most mammalian cells. Mammalian hepatocytes are an exception: they display transporting proteins on their sinusoidal surface (such as OATP1B1 and OATP1B3 of human hepatocytes [8-10]), which internalize phallotoxins and amatoxins. It is through the presence of the transporting protein OATP1B3 on human hepatocytes, for example, that amatoxins are feared as liver toxins. Similarly, amatoxins have been observed for most mammalian cells. This parameter was important, since cytotoxicity depends not only on membrane permeability but also on actin affinity. Moreover, this parameter will hint on whether phalloidin was cleaved from its carrier inside the cell, in cases where low actin affinity of a phallotoxin derivative was combined with high cytotoxicity. Relative affinity values of the phallotoxin derivatives to muscle actin are shown in Table 1.

The aim of this study was to explore known lipophilic and polycationic internalization-mediating moieties for their applicability with phallotoxins and amatoxins, and to find novel internalization-mediating moieties and investigate the potential of their conjugates with toxins as specific inhibitors. The internalization-mediating moieties used in this study were either lipophilic in nature (such as oleic acid), or multicationic (such as polylysine and octarginine): two features applying also either lipophilic in nature (such as oleic acid), or multicationic internalization-mediating moieties and investigate the potential of their conjugates with toxins as specific inhibitors. Thus, both the cytoskeleton and the eukaryotic transcription machine are complex structures and still under investigation, may explain the continuing interest in these two kinds of specific inhibitors.

Results

Preparation of phallotoxin derivatives

Attachment sites in phalloidin (Figure 1) for conjugation with uptake-mediating moieties were chosen based on our knowledge of structure–activity relationships in phalloidin [1]: As the side chains in the small heterodit peptide ring are known to be involved in actin binding they were not used for derivatization. In the larger peptide ring, on the other hand, the dihydroxylated leucine moiety is juxtaposed to the actin binding site and thus appeared as most promising for derivatization. Accordingly, all residues investigated in this study were attached to the C6-atom of the dihydroxylated leucine moiety, either as esters or amides (Table 1). Large residues, which might disturb the interaction of a phallotoxin derivative with actin through steric hindrance, were coupled via disulfide-containing linkers, which would be reduced inside the cell so as to release a defined thiol derivative of phalloidin (Table 1).

![Figure 1: Chemical structure of phalloidin with the attachment site (R) used for conjugation to uptake-mediating moieties.](image)

Actin binding

All phalloidin derivatives were tested for their affinity to muscle actin (\( \alpha \)-actin), which is used as a model for \( \beta \)-actin present in nonmuscle cells. This parameter was important, since cytotoxicity depends not only on membrane permeability but also on actin affinity. Moreover, this parameter will hint on whether phalloidin was cleaved from its carrier inside the cell, in cases where low actin affinity of a phallotoxin derivative was combined with high cytotoxicity. Relative affinity values of the phallotoxin derivatives to muscle actin are shown in Table 1.

Growth inhibition of mouse fibroblasts

By using the MTT cell proliferation assay, each phalloidin conjugate was examined for its capacity to inhibit the growth of mouse fibroblasts in vitro after 72 h incubation time (Table 1). Phalloidin displayed no antiproliferative activity up to a concentration of \( 1 \times 10^{-3} \) M in the culture medium. In contrast, the most lipophilic ester derivative, phalloidin oleate (1e), showed an IC\(_{50}\) value of proliferation inhibition of \( 2.5 \times 10^{-6} \) M, and was thus ca. 1000 times more active than phalloidin. Other esters (1a–1d) exhibited increasing cytotoxic activities with increasing hydrophobicity (Figure 2). In order to examine a possible relationship between cytotoxicity and hydrophobicity we determined the octanol/water equilibrium distribution coefficient of the ester derivatives (log \( P_{ow} \) values) and found a linear rela-
Table 1: Structures of phalloidin derivatives, their relative affinity values for α-actin as compared to phalloidin and their IC\textsubscript{50} values of cell growth inhibition after incubation for 72 h as determined by MTT cell proliferation assay (n.a.: not assayed).

<table>
<thead>
<tr>
<th>R</th>
<th>Name/designation</th>
<th>Relative affinity to α-actin [%]</th>
<th>IC\textsubscript{50} values NIH 3T3 fibroblasts [µM]</th>
<th>IC\textsubscript{50} values Jurkat cells [µM]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>OH phalloidin</td>
<td>100</td>
<td>ca. 1,000</td>
<td>ca. 1,000</td>
</tr>
<tr>
<td>1a</td>
<td>OCOC\textsubscript{6}H\textsubscript{5} (1)-benzoate</td>
<td>76</td>
<td>92</td>
<td>94</td>
</tr>
<tr>
<td>1b</td>
<td>OCOC\textsubscript{6}H\textsubscript{4}O-OH (1)-salicylate</td>
<td>81</td>
<td>364</td>
<td>96</td>
</tr>
<tr>
<td>1c</td>
<td>OCOC\textsubscript{7}H\textsubscript{15} (1)-octanoate</td>
<td>12</td>
<td>44</td>
<td>37</td>
</tr>
<tr>
<td>1d</td>
<td>OCOC\textsubscript{13}H\textsubscript{27} (1)-myristate</td>
<td>7</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td>1e</td>
<td>OCO(CH\textsubscript{2})\textsubscript{7}CHCH(CH\textsubscript{2})\textsubscript{7}CH\textsubscript{3} (1)-oleate</td>
<td>4</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>NH\textsubscript{2} aminophalloid</td>
<td>2</td>
<td>ca. 1,000</td>
<td>ca. 1,000</td>
</tr>
<tr>
<td>2a</td>
<td>NHCO\textsubscript{6}H\textsubscript{5} N-benzoyl-(2)</td>
<td>74</td>
<td>693</td>
<td>599</td>
</tr>
<tr>
<td>2b</td>
<td>NHCO(CH\textsubscript{2})\textsubscript{2}CHCH(CH\textsubscript{2})\textsubscript{3} N-oxy-ol(2)</td>
<td>5</td>
<td>883</td>
<td>630</td>
</tr>
<tr>
<td>2c</td>
<td>NHCO(CH\textsubscript{2})\textsubscript{2}SS(Ac)CysGlyTyrGly-Arg(Lys)\textsubscript{2}(Arg)\textsubscript{2}Glu(Arg)\textsubscript{2}OH (2)-Tat-peptide</td>
<td>3</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>2d</td>
<td>NHCO(CH\textsubscript{2})\textsubscript{2}SS(Ac)CysGly[Arg]\textsubscript{8}OH (2)-octarginine</td>
<td>2</td>
<td>11</td>
<td>7</td>
</tr>
<tr>
<td>2e</td>
<td>NHCO(CH\textsubscript{2})\textsubscript{2}CONH(Lys)\textsubscript{2}10</td>
<td>(2)-poly-(L)-lysine\textsubscript{28,000}</td>
<td>40</td>
<td>5</td>
</tr>
<tr>
<td>2f</td>
<td>NHCO(CH\textsubscript{2})\textsubscript{2}SS(CH\textsubscript{2})\textsubscript{2}CONH(Lys)\textsubscript{2}10</td>
<td>(2)-poly-(D)-lysine\textsubscript{28,000}</td>
<td>38</td>
<td>54</td>
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<tr>
<td>2g</td>
<td>NH(CH\textsubscript{2})\textsubscript{2}SS(CH\textsubscript{2})\textsubscript{2}CONH(PEG)\textsubscript{3000} (2)-SS PEG\textsubscript{800}</td>
<td>3</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>2h</td>
<td>NH(CH\textsubscript{2})\textsubscript{2}SS(CH\textsubscript{2})\textsubscript{2}CONH(PEG)\textsubscript{22,200} (2)-SS PEG\textsubscript{85,200}</td>
<td>4</td>
<td>77</td>
<td>50</td>
</tr>
<tr>
<td>2i</td>
<td>NH(CH\textsubscript{2})\textsubscript{2}SS(CH\textsubscript{2})\textsubscript{2}CONH(PEG)\textsubscript{23,000} (2)-SS PEG\textsubscript{23,000}</td>
<td>1</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>2f +DTT</td>
<td>NH(CH\textsubscript{2})\textsubscript{2}SH N-(2-mercaptoethyl)-(2)-SH</td>
<td>37</td>
<td>n.a.</td>
<td>n.a.</td>
</tr>
<tr>
<td>3</td>
<td>for structure see Figure 6a</td>
<td></td>
<td></td>
<td></td>
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</table>

Figure 2: Cytotoxicity of phalloidin derivatives. NIH 3T3 mouse fibroblasts were incubated with various concentrations of phalloidin and phalloidin derivatives. Cell viability was determined after 72 h incubation time by MTT assay.
Figure 3: n-Octanol/water distribution coefficients (log $P_{ow}$) of the hydrophobic phalloidin derivatives of Figure 2 plotted versus cell toxicity (log ($1/IC_{50}$)) in NIH 3T3 mouse fibroblasts.

Polycationic derivatives of phalloidin, such as the polylsine conjugates, were highly toxic for mouse fibroblasts, and their antiproliferative activity was comparable to the most lipophilic derivative, phalloidin oleate (Figure 4a). Their toxicity was found to be dependent on the configuration of the polymer, since phalloidin bound to D-configurated polylysine was about 10 times less toxic than when bound to poly-(L)-lysine. This suggests that the release of a toxic phalloidin species inside the cell includes the enzymatic breakdown of the carrier. In agreement with this we found no difference between the L-configurated and the D-configurated carrier when the linker contained a disulfide bridge, arguing for the presence of a disulfide-reducing compartment inside the cells.

Likewise, high cytotoxicity was found for the octarginine conjugate (Figure 4a). From the fact that the cytotoxic activities of the polylysine derivative (ca. 150 residues) and the oligoarginine derivative (8 residues) are comparable, we argue that arginine residues are more effective in mediating internalization of phalloidin than are lysine residues.

Figure 4: (a) and (b): NIH 3T3 mouse fibroblasts were incubated with various concentrations of phalloidin and phalloidin derivatives. Cell viability was determined after 72 h incubation time by MTT assay.
For phalloidin bound to methoxy-polyethylene-glycol we found that cytotoxicity strictly depended on the molecular weight of the polymer chain (Figure 4b). Most active was the conjugate with the longest chain ($M_r = 22,600$). Cytotoxicity fell strikingly when the polymer chain was shortened (Table 1).

Coupling to Tat-peptide was also very effective in enhancing the cytotoxicity of phalloidin (Figure 4b), while the phalloidin conjugate with the Kaposi protein fragment (likewise claimed to enhance membrane permeability) was much less effective (data not shown).

Beside mouse fibroblasts, we investigated the phalloidin derivatives in several human leukemia and lymphoma cell lines, in order to find possible specificities for one or the other kind of tumor cell. However, Jurkat cells (Table 1) and all other cell lines tested (K562 cells, HL-60 cells, and Daudi cells) showed sensitivities comparable to mouse fibroblasts (see Supporting Information File 1).

Uptake kinetics
Since the toxic effects of phalloidin develop slowly, growth inhibition was measured only after 72 h. During this period several partial processes must occur such as binding to the plasma membrane, internalization, processing and toxin release, etc., which cannot be distinguished. However, replacement of the toxin medium by toxin-free medium after various times of incubation would provide information on the time required for each of the toxin derivatives to bind to the cell surface. We compared the three most effective phalloidin conjugates, phalloidin oleate (1e), phalloidin-Tat conjugate (2c), and phalloidin polylysine (2e) (Figure 5) and found that the exposure times necessary to achieve, e.g., a 50% growth inhibition by a given toxin concentration after 72 h indeed varied considerably, from 2 h to 24 h. The results show that the polycationic derivative was bound much more rapidly than the lipophilic conjugate.

A fluorescent residue that enhances uptake of phalloidin into cells
Tetramethylrhodaminyl-phalloidin (Figure 6a) has been used to visualize actin fibers in fixed cells for 30 years. Here we show that the rhodamine residue also strongly enhanced cellular uptake, making this phalloidin derivative a tool for cell biology. With an IC$_{50}$ value of 11 µM its toxicity is comparable to those of the most toxic phalloidin derivatives, phalloidin oleate (1e) and phalloidin-poly-(L)-lysine (2e). More importantly, rhodaminyl-phalloidin seems not to be cleaved inside the cell and, through its fluorescence, can report on the structure of its target protein, the actin filaments, albeit under toxic conditions.

Using rhodamine-labeled phalloidin, we could also study how membrane-permeable peptides are incorporated into the cell. Immediately after exposure the toxin was located on the plasma membrane of the cells as shown by fluorescence microscopy (Figure 6c), while after 6 h increasing amounts of the toxin were found in endocytotic vesicles. After 24 h most of the rhodamine-labeled toxin was still in endosomes, while some of it had found its target, as concluded from the decoration of filaments.

Phalloidin causes apoptosis of cells
Under the microscope, cells treated with membrane-permeable phalloidin derivatives appeared shrunken and developed blebs, as described for cells undergoing apoptosis. Treatment with
Figure 6: (a) Chemical structure of tetramethylrhodaminyl-phalloidin (3). (b) Growth inhibition of NIH 3T3 mouse fibroblasts by tetramethylrhodaminyl-phalloidin (3) versus phalloidin (1) after 72 h incubation time. (c) Binding and uptake of tetramethylrhodaminyl-phalloidin (3) in NIH 3T3 mouse fibroblasts after 1, 6 and 24 h, as documented by fluorescence microscopy.

annexin followed by flow cytometric analysis showed a fluorescence distribution typical for apoptosis and similar to that induced by camptothecin. Cells treated with native phalloidin were indistinguishable from controls [13].

Growth inhibition by amatoxin derivatives

Unlike the phallotoxins, the natural amatoxins are toxic in cell cultures, exhibiting $IC_{50}$ values around $10^{-6}$ M. As amatoxins are more hydrophilic than phallotoxins, it seems unlikely that their membrane permeation capacity is larger than that of phallotoxins. The more likely explanation is that for amatoxins the threshold concentration lethal for cells is much lower than for phallotoxins (see Discussion).

Two of the internalization-mediating residues investigated in the phallotoxin series, oleic acid and polylysine, were also tested for their uptake capacity in the amatoxin series. Structure–activity studies had shown that the primary OH group of the dihydroxy-isoleucine moiety (Figure 7) is not involved in RNA polymerase II binding and, hence, may be used for derivatization with oleic acid chloride (Table 2). In order to avoid concurrent acylation of the 6’-OH of tryptophan, the phenolic OH was methylated before acetylation. For coupling to polylysine the natural carboxy group of aspartic acid as present in β-amanitin was used, which after activation as N-hydroxysuccinimide ester reacted with ε-amino groups in polylysine. Since

Figure 7: Chemical structure of amanitin with attachment site for conjugation to internalization-mediating moieties ($R^1$, $R^2$).
Table 2: Structures of amanitin derivatives and IC<sub>50</sub> concentrations of growth inhibition in mouse fibroblasts after 72 h incubation time (MTT cell proliferation assay).

<table>
<thead>
<tr>
<th>R&lt;sup&gt;1&lt;/sup&gt;</th>
<th>R&lt;sup&gt;2&lt;/sup&gt;</th>
<th>R&lt;sup&gt;3&lt;/sup&gt;</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt; values NIH 3T3 fibroblasts [nM]</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>OH</td>
<td>OH</td>
<td>α-amanitin</td>
</tr>
<tr>
<td>4a</td>
<td>OCO(CH&lt;sub&gt;2&lt;/sub&gt;)&lt;sub&gt;7&lt;/sub&gt;CHCH(CH&lt;sub&gt;2&lt;/sub&gt;)&lt;sub&gt;7&lt;/sub&gt;CH&lt;sub&gt;3&lt;/sub&gt;</td>
<td>OH</td>
<td>α-amanitin oleate</td>
</tr>
<tr>
<td>5</td>
<td>OH</td>
<td>OH</td>
<td>β-amanitin</td>
</tr>
<tr>
<td>5a</td>
<td>OH</td>
<td>OCH&lt;sub&gt;3&lt;/sub&gt;</td>
<td>β-amanitin poly-(L)-lysine</td>
</tr>
<tr>
<td>5b</td>
<td>OH</td>
<td>OCH&lt;sub&gt;3&lt;/sub&gt;</td>
<td>β-amanitin poly-(D)-lysine</td>
</tr>
</tbody>
</table>

Figure 8: NIH 3T3 mouse fibroblasts were incubated with various concentrations of α-amanitin and amanitin derivatives. Cell viability was determined after 72 h incubation time by MTT assay.

Discussion

Growth inhibition as a parameter of internalization

Phallotoxins as well as amatoxins find their targets inside the cell. Thus, if growth inhibition by these toxins occurs the toxins must have penetrated the cell. Moreover, the extent of the toxic lesions will mirror the amount of toxin that penetrated the cell and, hence, can be used to estimate the amount of toxin taken up. Although some of the steps involved in the uptake process became evident in this study it was not our aim to investigate

the polymeric carrier, as shown by the fact that β-amanitin coupled to the (L)-polymer was 1800 times more toxic than when coupled to the (D)-polymer (Table 2).
details of internalization, but to gather experience with moieties that may help to overcome the plasma membrane barrier and to shift hydrophilic peptides into a cell.

Previous experience with internalization-mediating moieties

Lipid acids have been used by various researchers to enhance the uptake of peptides and proteins into cells. For example Honeycutt et al. [14] used palmitic acid to deliver a protease inhibitor into cells, and Bradley et al. [15] used docosahexanoic acid to improve the uptake of paclitaxel into tumor cells. For a review see Wong and Toth [16]. Of particular interest was the incorporation of anti-sense oligonucleotides into cells by coupling them to lipophilic ligands as reported by Boutorin et al. [17], Letsinger et al. [18], and Shea et al. [19].

Polycationic carriers such as polylysine and polyarginine represent another, and possibly even more effective technique for delivering drugs into cells. Ryser and Shen [20] reported the internalization by polylysine of methotrexate and horse radish peroxidases; Leonetti et al. [21] the internalization of oligonucleotides; and Mulders et al. [22] the internalization of adenovirus into cells. A polylysine peptoid derivative was used by Murphy et al. [23] for “gene delivery”, and Emi et al. [24] used polyarginine as a carrier for gene transfer.

Protein transduction domains such as Tat-peptide, usually consisting of 10–30 amino acids, have been used to transport enzymes, drugs, liposomes and supermagnetic particles into cells [25-29]. In the meantime it has been shown that nona-arginine is many times more efficient than Tat-peptide, suggesting that the internalization effect of Tat-peptide is mainly due to its eight cationic side chains [30].

Attachment to plasma membrane determines the rate of internalization

Plasma membranes of cells in culture represent a lipophilic phase in an aqueous medium, and amphiphilic compounds brought into this system will distribute between the two phases according to their nature. This process is similar to the distribution the substance will take between n-octanol and water, an idea that has been suggested by Palm et al. [31].

Compounds 1a–1e represent a series of phalloidin derivatives with increasing log $P_{ow}$ values. At the same time they represent a series of phalloidin derivatives with increasing cytotoxic activity. Since the two parameters are linearly related (Figure 3), we conclude that the amount of toxin attached to the membrane determines the extent of the toxic effects observed. Clearly, this conclusion is valid only under the condition that, as in our case, all derivatives belong to the same chemical class, esters for example, and that processing inside the cell will yield the same toxic product, here native phalloidin, set free in all cases.

Attachment to the plasma membrane can also occur by electrostatic forces and, thus, may be behind the internalization effect observed with polylysine or polyarginine as well. Plasma membranes expose numerous negatively charged components that can attract oligo- or polycationic molecules. As already pointed out, arginine residues as in 2d are more effective than lysine residues in 2e, as 120 lysine groups are required to match the internalization increase caused by eight arginine residues.

Concerning Tat-peptide, we believe that its eight cationic side chains interacting with the plasma membrane represent the more likely explanation for its internalization than other models proposed in the past.

Finally, tight attachment to the plasma membrane may also explain why the methoxy-polyethylene glycol residue mediates internalization, since polyethylene glycols are soluble not only in water but also in diethylether, i.e., can adopt a conformation capable of anchoring to the phospholipids in the plasma membrane. Lipophilic interaction of the aromatic part of tetramethylrhodamine with the plasma membrane is also thought to be the main cause of the internalization capacity of this fluorescent residue, beside its delocalized cationic charge on the two nitrogen atoms.

Although proved for a lipophilic phalloidin derivative only, we postulate that attachment to the plasma membrane also provides the first step in entering cells for the polycationic and the peglated toxin derivatives presented in this study.

Internalization and processing

The uptake process following the binding step of the phalloidin derivative 3 to the plasma membrane was identified as endocytosis. As shown in Figure 6c, the fluorescent phalloidin is bound on the plasma membrane of mouse fibroblasts after 1 h, while after 6 h most of the fluorescent material resides in endocytotic vesicles. This finding confirms the earlier observation [6] that TRITC-phalloidin 3 enters isolated mouse hepatocytes by endocytosis and not through a phalloidin-transporting protein, such as the OATP1B1 present on human hepatocytes. The type of endocytosis seen here remains, however, to be elucidated. Likewise, we have no data on whether lipophilic or polycationic phalloxins enter cells in the same way as the fluorescent toxin, but this seems likely since proteolytic enzymes present in endosomal-lysosomal compartments were found to be involved in the processing, and the difference in de-
rhodamine-labeled phalloidin seems useful as a tool for
derivative still decorates filaments in the cell. This property of
phalloidin, by a factor of >100. Linked by a thiourea moiety, as
ethylrhodaminyl residue can facilitate internalization, e.g., of
Likewise new, is the observation that the red fluorescent tetram-
active form. That after internalization the drug is released in a defined and
2i
carrier finds its way out of the endosomal-lysosomal
compartments into cytoplasm, where the target, the actin fila-
ments, is located.

Conclusions
With IC_{50} values of >10 mM (8 mg/mL; M_r 789 g/mol), phal-
loidin has so far been of no benefit for research on living cells,
unless it was microinjected. This is regrettable, since the molec-
ular mechanism of phalloidin action, the immobilization of the
microfilament system, has been investigated in great detail.
Moreover, phalloidin action is comparable to that of taxol,
which induces comparable immobilization of the microtubular
system and has been widely employed in cell research and even
in tumor therapy [32,33]. One difference is that taxol is a
lipophilic compound, and is thus active on cells at much lower
concentrations. We balanced this disadvantage by coupling phalloidin, e.g., to oleic acid, allowing studies of the molecular
of phalloidin on cells in the micromolar range and
encouraging its use in tumor therapy.

The internalization-mediating effect observed with phalloidin
was seen also with amatoxins. Although amanitin is active on
cells in the micromolar range, it may be advantageous to use it
bound to oleic acid or polylysine and thus decrease the medium
concentration necessary to achieve growth inhibition down to
the nanomolar range. Amanitin as a drug bound to tumor mono-
clonal antibodies, has recently been described for the therapy of
adenocarcinomas [34].

Coupling to methoxypolyethyleneglycol has been described as a
method to improve the solubility of drugs, prolong their half-
lives in plasma, or modulate their pharmacokinetics [35]. An
effect not described to our knowledge so far is that pegylation
can enhance, several hundred-fold, the penetration of a
hydrophilic drug into cells. Linkers with the drug should
contain a disulfide bridge, as in compound (2i), to make sure
that after internalization the drug is released in a defined and
active form.

Likewise new, is the observation that the red fluorescent tetrarn-
ethylrhodaminyl residue can facilitate internalization, e.g., of
phalloidin, by a factor of >100. Linked by a thiourea moiety, as
in compound 3, the fluorescent moiety is not cleaved inside the
cell, as concluded from the observation that this phalloidin
derivative still decorates filaments in the cell. This property of
rhodamine-labeled phalloidin seems useful as a tool for
studying the kinetics of phalloidin-induced disturbances in the
actin system of a cell in correlation with, for example, the
development of apoptosis in the cell.

Experimental
Fatty acid esters of phalloidin
Ten micromoles of phalloidin was dissolved in 0.1 mL dry pyri-
dine and reacted with 0.3 µmol of benzoyl chloride, salicyl
chloride, octoyl chloride, myristoyl chloride, or oleoyl chloride
for 2 h at rt. Under these conditions, the reaction proceeded
predominantly at the primary OH group of the γ,δ-dihydroxy-
leucine residue in position 7 of phalloidin, with ca. 10–20% of
esterification at the secondary OH groups. The reaction was
stopped with 2 mL of methanol, and solvents were removed in
vacuo at 60 °C. Purification of the esters was achieved by
preparative TLC on silica (Merck HF_254, Darmstadt; Germany)
in chloroform/methanol/2 N acetic acid (65:25:4), followed by
further purification of the methodanolic extract of the silica on a
Sephadex-LH20 column developed with methanol. Yields of
the esters were 43% for benzoyl phalloidin, 35% for salicyl
phalloidin, 46% for octoyl phalloidin, 50% for myristoyl phal-
loidin and 44% for oleoyl phalloidin; purities were 88 to 95% by
HPLC.

Aminophalloidin
Ten micromoles of toluene-4-sulfonyl chloride in 2 mL chloro-
form were added to 0.6 µmol of phalloidin in 5 mL dry pyri-
dine and allowed to react for 30 min on ice. The reaction was
stopped by the addition of 50 mL of dry diethyl ether, and the
sediment was isolated by centrifugation, washed twice with
50 mL diethyl ether, and dissolved in 5 mL methanol for separ-
ation on a Sephadex-LH20 (60 × 3 cm) column developed with
methanol/H_2O (1:1). Yield of the monotosylphalloidin was
65%; purity 94%. The vacuum-dried monotosylphalloidin was
dissolved in 40 mL of methanol containing 2.5 N ammonia and
reacted for 2 h. After evaporation in vacuo at 60 °C, the
aminophalloidin was purified on a Sephadex-LH20 column
with methanol as eluant. Yield of aminophalloidin was 80%,
purity 89% by HPLC.

Fatty acid amides of phalloidin
Aminophalloidin and the fatty acid chlorides were reacted as
described for the synthesis of phalloidin esters.

Linear peptides linked to aminophalloidin
Linear peptides such as Ac-Cys-Gly-Tyr-Gly-Arg-Lys-Lys-
Arg-Arg-Gln-Arg-Arg-Arg-OH (Tat peptide), and Ac-Cys-Gly-
Arg-GluOH (ArgO) were synthesized on an Eppendorf Ecosyn P
solid-phase synthesizer by using 9-fluorenylmethoxycarbonyl
(Fmoc)-Arg(Pbf)SPHB resin (Rapp Polymere, Tübingen,
Germany). For the Kaposi sequence Ac-Cys-Gly-Ala-Ala-Val-

Ala-Leu-Leu-Pro-Ala-Val-Leu-Leu-Ala-Leu-Ala-Pro-OH an Fmoc-Pro-Trt-Tentagel resin was used. All amino acids were incorporated with the α-amino functions protected with the Fmoc group. Side chain functions were protected as tert-butyl ethers (tyrosine), tert-butyl oxy carbonyl derivatives (lysine), trityl derivatives (cysteine, glutamine), and as (2,2,4,6,7-pentamethyl)dihydrobenzofuran-5-sulfonyl derivative (arginine). Coupling was performed by using a 4-fold excess of each of the protected amino acids and the coupling reagent 2-(1H-benzo triazole-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate (TBTU) and 2 equiv of diisopropyl ethylamine (DIEA) over the resin loading. Before the coupling steps the Fmoc groups were removed from the last amino acid of the growing peptide fragment by using 25% piperidine in dimethylformamide. After cleavage of the N-terminal Fmoc group, the peptide was removed from the resin under simultaneous cleavage of the amino-side-chain protecting groups by incubation for 3 h in a mixture of trifluoroacetic acid (TFA) (12 mL), ethanedithiol (0.6 mL), anisole (0.3 mL), water (0.3 mL), and trisopropyl silane (0.15 mL). The mixture was filtered, and washed with TFA and anhydrous diethyl ether. The crude products were further purified by HPLC on a Nucleosil 100 C18 (7 µm) column (250 × 10 mm, Macherey & Nagel, Düren, Germany) by using a gradient from 10–90% B in 32 min (solution A: 0.07% TFA/H2O; solution B: 0.059% TFA in 80% acetonitrile). The elution was monitored at 214 nm. The peptides were assayed for purity by analytical HPLC and ESI-MS. The peptides were coupled to aminophalloidin through the hetero-disulfide group in aminophalloidin to form an amide linkage. In general, the activated bifunctional cross-linking reagents DSP (dithiobis(succinimidyldioxybutyrate); Lomant’s reagent) with cleavable disulfide group, or DSS (disuccinimidyl suberate) containing a hydroxysuccinimide ester). Thus, the activated ester end of excess SPDP was reacted with the primary amine group in aminophalloidin to form an amide linkage. In general, 63 µmol aminophalloidin were dissolved in 3.4 mL H2O and 1.7 equiv SPDP, dissolved in 900 µL dimethylformamide, were added. The solution was adjusted to pH 7.5 with 1 N NaOH; reaction time was 1 h. Separation of the products was performed by Sephadex-LH20 column developed with methanol. The purity of PDP-aminophalloidin was 93%; the yield was about 65%. The 2-pyridyldithio group at the other end of the linker was reacted with the sulffhydryl in the amino terminal Ac-Cys-Gly moiety of the linear peptides to form a disulfide group. The reaction conditions of the three linear peptides varied slightly and were as follows: 33 µmol Tat-peptide, dissolved in 1.0 mL PBS was added to 40 µmol of PDP-aminophalloidin dissolved in 0.5 mL methanol; 13.6 µmol Arg8-peptide, dissolved in 1.5 mL PBS was added to 28 µmol of PDP-aminophalloidin dissolved in 0.25 mL methanol; 12 µmol Kaposi-peptide, dissolved in 0.8 mL PBS was added to 36 µmol of PDP-aminophalloidin dissolved in 0.2 mL methanol. Reaction time was in all cases 16 h at rt. Separation was achieved on Sephadex-LH20 with H2O/methanol (4:1) as solvent. Yield for Tat-phalloidin was 52%, for Arg8-phalloidin 43% and for Kaposis-peptide 39%. Purity of all conjugates was >90% as shown by HPLC (Table 1) and MALDI–TOF analysis. The HPLC analysis conditions were as follows: Column Knauer RP Nucleosil-100 C18 (250 × 4 mm); mobile phase was a linear gradient of buffer A H2O, 0.05% TFA and buffer B acetonitrile/H2O (9:1), 0.05% TFA.; flow rate, 1.2 mL/min.

General MALDI–TOF analysis was performed in the linear, high-mass, positive-ion mode with pulsed (time-delayed) extraction on a Kratos Maldi IV instrument (Shimadzu Deutschland, Düssiburg, Germany). Samples (usually in 0.1% TFA) were either applied by a sandwich technique (in which 0.7 µL of matrix was dried onto the sample spot, followed by 0.7 µL of sample and then another 0.7 µL of matrix) or, for more concentrated samples, were simply mixed 1:10 with the matrix solution and 0.7 µL of this mixture was dried onto the stainless-steel sample holder. Matrix solutions were usually α-cyano-4-hydroxycinnamic acid [dissolved at 10 mg/mL in 50% acetonitrile, 50% 0.1% TFA (all % v/v)]. Spectra were calibrated by using near-external standards consisting of a mixture of fragment 1–4 of substance P ([M + H] m/z 497.6); angiotensin II ([M + H] m/z 1047.2); angiotensin I ([M + H] m/z 1297.5); fragment 1–13 of angiotensinogen ([M + H] m/z 1646.9); and oxidized insulin B chain ([M + H] m/z 3496.9). Absolute m/z values occasionally varied up to 1 Da depending on the individual calibration and the distance and time between the individual measurements; for these samples, spectra were recalibrated by using known m/z values of the largest peak(s). Spectra were collected and analyzed by using standard Kratos software (Sun OS, Release 5.4, OpenWindows Ver. 3.4, Kratos Kompackt Software Ver. 5.2.0) and were usually the average of 50–100 individual laser shots across the width of the sample spot. Data were smoothed and baseline-corrected, generally with a window width of 30 channels.

Polymers linked to aminophalloidin

Poly-(L)-lysine (hydro bromide; Mw = 27500), and monom ethoxy-polyethyleneglycolamide (Mr = 810, 5200, 22600) were coupled to aminophalloidin by the amine-reactive homobifunctional cross-linking reagents DSP (dithiobis(succinimidylpropionate); Lomant’s reagent) with cleavable disulfide group, or DSS (disuccinimidyl suberate) containing a hydrocarbon chain instead of the disulfide group. DSP or DSS (248 µmol) were dissolved in 1.0 mL of N,N-dimethylformamide and added to 63 µmol dried aminophalloidin. The reaction was started with 2 µL triethylamine and allowed to proceed under magnetic stirring for 16 h at rt. The reaction was stopped with 10 mL diethyl ether and the mixture was centrifuged; after a second wash with ether, the sediment was dissolved in 5 mL.
methanol and separated on a Sephadex-LH20 column with methanol as solvent. Yield of DSP-, and DSS-phallolidin was about 80%, purity >90% for both.

Seven milligrams DSP-, or DSS-phallolidin were dissolved in 0.5 M N,N-dimethylformamide and 5 equiv of poly-(L)-lysine hydrobromide or poly-(D)-lysine hydrobromide added in 0.5 mL PBS. After reaction for 16 h at rt, high-molecular-weight products were separated by gel-filtration chromatography with Sephadex G-25 with 0.1% NaCl as eluant. After lyophilisation, the amount of phallolidin coupled to the polymer was determined from the characteristic absorption of phallolidin at 300 nm (ε = 10,100). We found that ca. 1 out of ca.10 lysine residues was spiked with aminoaphallolidin, independent of the molecular weight of the polymer. For modification of DSP-phallolidin and DSS-phallolidin with methoxypolyethylene glycol-lamine, monomethoxypolyethylene glycol was tosylated and reacted with ammonia to yield monomethoxy-PEG with a reactive amino group: 100 mg monomethoxy-PEG 810, 5,200 and 22,600 were dissolved in 1.0 mL of dry pyridine in a round-bottom flask on ice, and 5 equiv tolul-4-sulfonylchloride in 0.4 M chloroform were added dropwise. After being stirred for 30 min, the reaction was stopped with 20 mL of diethyl ether. Sediment was dried in a rotation evaporator and reacted with 20 mL of methanol/2.5 N ammonia. After 1 h, the solution was evaporated in vacuo and the aminomonomethoxy-PEG purified by Sephadex-LH20 chromatography. Ten milligrams DSP- or DSS-a minophallolidin were dissolved in 0.5 mL N,N-dimethylformamide and added to 1 equiv dry aminomonomethoxy-PEG. After reaction for 16 h at rt, phallolidin PEG 22,600 and phallolidin PEG 5,200 were purified on a Sephadex G25 column by using 0.1% NaCl as solvent, and phallolidin PEG 810 was purified on a Sephadex-LH20 column developed with methanol. The yield was 37% for phallolidin PEG 22,600, 34% for phallolidin PEG 5,200 and 45% for phallolidin PEG 810.

Affinity to rabbit muscle actin
Actin was prepared from rabbit muscle as described previously [36]. The binding assay was used with the following modifications: Freshly prepared G-actin solution was diluted in Tris-ATP buffer (2 mM of Tris; 0.2 mM ATP; 0.1 mM CaCl2; 0.02% NaCl; pH 7.8) to an extinction of 0.28 at 290 nm (null balance at 310 nm). Typically, 18 µL of a [3H]-demethylphalloin methanolic solution (specific activity 9 Ci/mmol) was evaporated in vacuo and the aminomonomethoxy-PEG purified on a Sephadex-G25 column with 0.1% NaCl as eluant. After lyophilisation, the amount of phallolidin coupled to the polymer was determined from the characteristic absorption of phallolidin at 300 nm (ε = 10,100). We found that ca. 1 out of ca.10 lysine residues was spiked with aminoaphallolidin, independent of the molecular weight of the polymer. For modification of DSP-phallolidin and DSS-phallolidin with methoxypolyethylene glycol-lamine, monomethoxypolyethylene glycol was tosylated and reacted with ammonia to yield monomethoxy-PEG with a reactive amino group: 100 mg monomethoxy-PEG 810, 5,200 and 22,600 were dissolved in 1.0 mL of dry pyridine in a round-bottom flask on ice, and 5 equiv tolul-4-sulfonylchloride in 0.4 M chloroform were added dropwise. After being stirred for 30 min, the reaction was stopped with 20 mL of diethyl ether. Sediment was dried in a rotation evaporator and reacted with 20 mL of methanol/2.5 N ammonia. After 1 h, the solution was evaporated in vacuo and the aminomonomethoxy-PEG purified by Sephadex-LH20 chromatography. Ten milligrams DSP- or DSS-aminophallolidin were dissolved in 0.5 mL N,N-dimethylformamide and added to 1 equiv dry aminomonomethoxy-PEG. After reaction for 16 h at rt, phallolidin PEG 22,600 and phallolidin PEG 5,200 were purified on a Sephadex G25 column by using 0.1% NaCl as solvent, and phallolidin PEG 810 was purified on a Sephadex-LH20 column developed with methanol. The yield was 37% for phallolidin PEG 22,600, 34% for phallolidin PEG 5,200 and 45% for phallolidin PEG 810.

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MTT proliferation assay
Mouse fibroblasts NIH 3T3 (generous donation from Prof. Traub, MPI Ladenburg), K562, HL-60 and Jurkat cells (ATCC) maintained in RPMI 1640 medium containing 10% fetal calf serum and 0.05 mM β-mercaptoethanol, were cultured at 37 °C in a humidified 95% air/5% CO2 incubator. The cytotoxicity of phallotoxins was assessed by using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyloxazolium bromide (MTT) assay. Exponentially growing cells were plated at a density of 2 × 104 cells/well in 96-well plates 24 h before the toxin was added in medium with up to 1% DMSO, with a volume equal to the volume of medium in the culture dish. The final concentrations of toxins in the media were between 10−3 and 10−9 M. At 72 h, the medium was replaced by serum-free medium containing 25 µL MTT solution (5 mg/mL in PBS), and the incubation was continued at 37 °C for 4 h. Then, lysis buffer (100 µL 20% SDS in 50% dimethylformamide) was added to each well and incubated for another 16–20 h. Viability of cells was determined by measuring the 570 nm absorbance of each well using a microplate reader (Molecular Devices). IC50 values were calculated as the concentration of toxin required to reduce the absorbance to 50% of the control cultures.

Microscopic studies
Exponentially growing fibroblasts were plated at a density of 2 × 104 cells/well in glass-bottom dishes 24 h before the fluorescently labeled peptides were added. The final concentration of the peptides was 10−5 M. After different incubation times the cells were washed with fresh medium, and microscopic studies were performed by using a confocal laser scanning microscope TCS SP2 (Leica Microsystems, Heidelberg/Mannheim, Germany), equipped with an inverted microscope DMI2E2 and an incubation chamber (Pe-Con Erbach, Germany). Image data stacks and time-lapse studies of the live cells were obtained at 37 °C and in 5% CO2 atmosphere with a 100×/1.4 N.A.
oil immersion objective and CLSM software (Leica Microsystems). The data were processed with ImageJ software, optimizing images in brightness and contrast, and visualization of the fluorescent structures was performed on serial confocal optical sections.

Supporting Information

Supporting Information File 1
Structures of phalloidin derivatives and IC_{50} concentration values of cell growth inhibition.

References

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