Oligonucleotide Phosphorothioates Enter Cells by Thiol-Mediated Uptake

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Abstract: Oligonucleotide phosphorothioates (OPS) are DNA or RNA mimics where one phosphate oxygen is replaced by a sulfur atom. They have been shown to enter mammalian cells much more efficiently than non-modified DNA. Thus, solving one of the key challenges with oligonucleotide technology, OPS became very useful in practice, with several FDA-approved drugs on the market or in late clinical trials. However, the mechanism accounting for this facile cellular uptake is unknown. Here, we show that OPS enter cells by thiol-mediated uptake. The transient adaptive network produced by dynamic covalent pseudo-disulfide exchange is characterized in action. Inhibitors with nanomolar efficiency are provided, together with activators that reduce endosomal capture for efficient delivery of OPS into the cytosol, the site of action.

Of general significance for science and society in the broadest sense, oligonucleotide technology applications have often been hampered by poor cellular uptake.\[1–3\] Besides a large number of gene transfection vectors,\[1–3\] many non-native modifications of oligonucleotides have been introduced in the past decades to address this challenge.\[4\] Oligonucleotide phosphorothioates (OPS) such as 1, where one oxygen atom of the bridging phosphodiester of the biological original 2 is replaced by a sulfur atom, have been one of the first backbone modification introduced in the field (Figure 1).\[5\] Apart from better nuclease stability and hydrophobicity, OPS have been shown to penetrate cells much more efficiently than unmodified DNA in the absence of transfecting agents (Figure 1 A, B).\[6\] For these reasons, OPS have found applications in the past decades in the clinics, with several FDA-approved drugs on the market and several in late clinical trials.\[7\]

Considering a pKₐ close to zero for phosphorothioate monomers,\[8\] transient protonation for transmembrane translocation in neutral form\[9\] was not likely to account for cell penetration. Pioneering work by Crooke and co-workers has shown that interactions with membrane-bound proteins are essential for OPS uptake through endocytosis and other, unclear, pathways.\[7,10\] Among identified protein partners, many of them are disulfide-rich proteins.\[7,10–15\] This supported that dynamic covalent exchange chemistry\[11,13,16,17\] might enable OPS to penetrate cells so easily. Thiol-mediated uptake is emerging as method of choice to bring challenging substrates into cells and to hinder viral entry.\[1,10,17–19\] In the following, we report that thiol-mediated uptake accounts for the entry of OPS, and show how this knowledge is of use to enhance uptake and reduce endosomal capture.

The occurrence of thiol-mediated uptake is most convincingly demonstrated by the inhibition of the dynamic covalent exchange with the cell during uptake.\[10\] Traditionally, this has been done with Ellman’s reagent. Weak and unreliable, this single inhibitor has been replaced recently by a collection of inhibitors which are up to 5000 times more active and cover at least some of the different uptake pathways involved.\[20\]

From this collection, inhibitor candidates 3–12 were selected to explore interference with the uptake of OPS 1, a random 18-mer labelled with Cy5 at its 5'-terminus (Figure 1, Figure 2; Table S1).\[20\] If needed, inhibitors were prepared by multistep synthesis following reported procedures (see the Supporting Information). A recent automated high-content high-throughput (HCTH) screening assay\[20\] was

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Figure 1. CLSM images of HeLa Kyoto cells after 2 h incubation with DNA 2 (A) and OPS 1 without (B) and with (C) preincubation with 3 (10 μM); red, 1, 2 (Cy5); blue, Hoechst 33342; scale bar: 50 μm. R = nucleobases, sequence: AGCTCCCATACACCGAC.
used to secure data from multiwell plates with thousands of HeLa Kyoto cells. They were first preincubated for 1 h with 3–12 at various concentrations below their toxicity limit[20] and then, after inhibitor removal, incubated for 2 h with Cy5-OPS 1. The dependence of the uptake of 1 on the concentration of 3–12 (Figure 2A) was then used to determine the MIC \( \text{OPS} \) (Figure 2B) and, if accessible, also the IC\(_50\), i.e., the concentration to observe \( \approx 15\% \) and 50\% inhibition, respectively (Figures 1, 2, S10–S19, Table S2).

The best inhibitor of the uptake of \( \text{OPS} \) 1 was the hypervalent iodine reagent 4, which irreversibly reacts with thiols on the cell surface.[21] A sub-micromolar MIC \( \text{OPS} \) = 0.3 \( \mu \text{M} \) and an IC\(_{50}\) = 14 \( \mu \text{M} \) were obtained. Excellent inhibition of \( \text{OPS} \) uptake was also found for the most reactive cyclic oligochalcogenides (COCs), i.e., BPS pentasulfide 5.[22] ETP tetrasulfide 3[22] and ETP disulfide 6.[23] The less reactive AspA\((10,12)\) 9 was also less impressive as inhibitor. The cyclic thiosulfonate COC 8 was of interest mostly because, among other activities,[14] it best inhibited the cellular entry of SARS-CoV-2 virus.[26,27] Thus, 5′-AMPS 13 was used as mimimalist \( \text{OPS} \) to explore the possible exchange with the non-activated disulfides 14 of cystine residues on cell surfaces. The in situ formation of pseudo-disulfides 15 and their reduction with TCEP could be demonstrated by HPLC and MS (Figures 3B, S31). In contrast, 5′-AMPS 13 did not react with reduced cysteines (Figure S32).

These results supported that the dynamic covalent chemistry of \( \text{OPS} \) 1 operates with non-activated disulfides but not with thiols (Figure 3, II). The efficient inhibition with thiol-reactive agents thus implied that the blocking of exofacial thiols also led to the blocking of exofacial disulfides via biological regulation by, e.g., PDI (protein disulfide isomerase) or glutathione, to ultimately inactivate all accessible sulfur, thiols and disulfides (V).[10,14,28] Alternatively, or in addition, our inhibition results could imply that \( \text{OPS} \) 1 is dynamic covalently activated by extracellular disulfides near the cell surface, e.g., oxidized glutathione, and the resulting, transient pseudo-disulfides 1 then exchange with exofacial thiols (IV) to end up with the same exchange cascade (III).

The possibility to activate \( \text{OPS} \) 1 in situ as pseudo-oligochalcogenides 1 was intriguing (Figure 3A). The same inhibitors identified above could conceivably act as activators.

Dependent on the transferrin receptor, while ETP 6 is not[12,23]. Compared to ETP 6a, uptake inhibition of \( \text{OPS} \) 1 was better with irreversible 4 and AspA 9 but weaker with ETP 6. This could support pathways including the transferrin receptor, which would agree with significant endosomal capture (vide infra).[10] High activity of ETP 3 was intriguing because, contrary to the contracted 6, the expanded 3 is a poor transporter,[22] Ebselen analog 11, an inhibitor of the entry of 6a (unpublished) and SARS-CoV-2,[24] was completely inactive up to its solubility limit. In contrast, the anomalous dose response of Ellman’s reagent 10 against 6a[20] converted into normal curves with \( \text{OPS} \) 1, which provided also access to an IC\(_{50}\) = 3.6 \( \mu \text{M} \).

The overall positive correlation in inhibition patterns of \( \text{OPS} \) and ETP 6a uptake (Figure 2B) might imply that the many proteins known from \( \text{OPS} \) uptake[17] contribute to the entry of COCs and viruses. SCARB1, for example, is involved in the uptake of \( \text{OPS} \),[20] SARS-CoV-2,[11] hepatitis virus,[10] and COC 9.[12] Hinting toward thiol-mediated uptake as a unifying network coding for entry, similar, at least partial coincidences can be found for other target candidates (e.g. the transferrin receptor,[10,12] EGFR,[7,10,14] integrins,[7,25] and CLIC[7,12]).

Thiol-mediated uptake is defined as enhanced, inhibitable uptake in the presence of dynamic covalent chalcogen exchangers, usually disulfides.[10] This cascade exchange chemistry occurs during uptake via direct translocation, fusion or endocytosis, involves multiple protein targets (vide supra) and, presumably, also transient micellar membrane microdomains, at least for the cytosolic delivery of large substrates.[10] In this context, the inhibition of \( \text{OPS} \) uptake by thio-reactive probes was intriguing because the multivalent \( \text{OPS} \) 1 would be expected to exchange with disulfides rather than with thiols on the cell surface (Figure 3A, II) to trigger the exchange cascades expected for efficient uptake (III).

Surprisingly little is known about the dynamic covalent exchange chemistry of phosphorothioates.[26,27] Thus, 5′-AMPS 13 was used as minimalist \( \text{OPS} \) to explore the possible exchange with the non-activated disulfides 14 of cystine residues on cell surfaces. The in situ formation of pseudo-disulfides 15 and their reduction with TCEP could be demonstrated by HPLC and MS (Figures 3B, S31). In contrast, 5′-AMPS 13 did not react with reduced cysteines (Figure S32).

Comparison with results from the inhibition of the thiol-mediated uptake of fluorescent ETP 6a revealed an overall positive correlation of MIC\(_{\text{OPS}}\) and MIC\(_{\text{ETP}}\) consistent with their reactivities, but with some distinct outliers (Figure 2B). This mismatch has been observed previously comparing 6a and 5a and interpreted as support for the existence of multiple targets.[25] Multiple pathways available for thiol-mediated uptake were implied also from proteomics analysis, which revealed, inter alia, that the entry of AspA 9 is...
Figure 3. A) A tentative dynamic covalent exchange mechanism for the thiol-mediated uptake of OPS. B) Normalized HPLC traces for the exchange of 13 (A = adenosyl) with 14a (top to bottom: 14a, 1:1, 10:1, 100:1) 13/14a and 100:1 after addition of 100 equiv. of TCEP, and mass spectrum of 15b. Conditions: 100 μM 14a, PBS, pH 7.4; *=−reduced 14a.

Depending on conditions. While their incubation with cells removes cell surface thiols and thus inhibits thiol-mediated uptake (Figure 3 A, V), the complementary incubation with OPS 1 could possibly afford pseudo-oligochalcogenides to facilitate exchange with cell surface thiols and thus activate thiol-mediated uptake (Figure 3A, I). The exchange of 5'-AMPs 13 with DTNB 10 to produce activated pseudo-disulfides was confirmed easily (Figure S29). MMTS 12 was found to exchange most efficiently with 13, with only 4 equivalents needed to reach full conversion into pseudo-disulfide 16 (Figures 4G, S30). Exchange of phosphorothioate monomer 17 with BPS 5a triggered the emergence of an adaptive dynamic polysulfide network [26] on the OPS model 18 (Figure 4H). With the same HPLC-MS fingerprinting, transient, dynamic covalent exchange activation was confirmed also for activators 3 and 7 (Figures S27, S28).

To activate in situ for thiol-mediated uptake, OPS 1 was incubated for 30 min with various concentrations of potential activators (Figures S20–S26). Added as mixtures, the uptake of OPS increased with increasing concentrations of activators up to concentrations close to the MIC, beyond which the leftover activator started to inhibit uptake (e.g., Figure 4D, S20, S21, S23). The removal of the excess activator was possible but not always beneficial, since this led to a partial loss of activation due to the transient nature of activated OPS 1, as confirmed by HPLC (e.g., Figure S28). Thus, transient activation of OPS 1 was examined without purification, which limited activation to concentrations below the onset of overcompeting inhibition (Figure 4D). With ETP 3, BPS 5, polysulfides 7, and thiosulfonates 8 and 12, activation remained partially preserved after removal of excess activator, resulting in up to ≈1.7-fold increase with 3 (Figures 4A–C, S20).

Like the transferrin-receptor dependent AspA 9, OPS are known to enter cells mostly by endocytosis, localizing in early and late endosomes within 10–50 min and in lysosomes afterwards, with activities observed only after several hours, indicating that endosomal escape is slow. After OPS activation, more diffuse fluorescence was observed in CLSM images (Figure 4A, B). Image analysis was performed...
to segment punctate structures that, with all likelihood, correspond to endolysosomes. Whereas the integrated intensity in the whole cell increased upon activation, the integrated fluorescence intensity remained constant in the mask (Figure 4E). This difference indicated that the increase of uptake is correlated with a more diffuse, most likely cytosolic localization. The same shift from endosomal to cytosolic localization. The same shift from endosomal to cytosolic fluorescence intensity remained constant in the mask (Figure 3A). To close, we reiterate that HCHT imaging automatically informs on cell viability, and that all reported data were obtained at concentrations below the onset of toxicity (which already has been reported for most inhibitors used).

In summary, the biology of OPS is understood and not topic of this study. It is also known that the success of OPS in biology and medicine originates in part from their ability to penetrate cells, and several proteins have been identified to contribute to endocytosis and other, unknown mechanisms. What has remained mysterious is the question why the replacement of one oxygen by one sulfur per monomer in the backbone converts an oligonucleotide that cannot penetrate cells into one that can. Here we show that the underrecognized dynamic covalent exchange of phosphorothioates with cellular thiois and disulfides accounts for the cell penetration, and that their thiol-mediated uptake can be inhibited and activated according to the general principles of dynamic covalent sulfur exchange chemistry, from simple pseudodisulfides over thiosulfonates to more complex adaptive networks.

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Conflict of Interest

The authors declare no conflict of interest.

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