Enhancement of the Click Chemistry for the Inverse Diels Alder Technology by Functionalization of Amide-Based Monomers

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Abstract

In the near future personalized medicine with nucleic acids will play a key role in molecular diagnostics and therapy, which require new properties of the nucleic acids, like stability against enzymatic degradation. Here we demonstrate that the replacement of nucleobases with PNA by functional molecules harbouring either a dienophile or a diene reactivity is feasible and confers all new options for functionalization. These newly developed derivatives allow independent multi-ligations of multi-faceted components by use of the inverse Diels Alder technology. The high chemical stability and the ease of synthesis qualify these polyamide building blocks as favourites for intracellular delivery and targeting applications. This allows local drug concentrations sufficient for imaging and therapy and simultaneously a reduction of the application doses. It is important to point out that this technology is not restricted to ligation of medicament material; it is also a candidate to develop new and highly efficient active compounds for a “sustainable pharmacy”.

Key words: Click Chemistry; Diels Alder Reaction inverse (DARinv); local concentration; Peptide Nucleic Acid (PNA); PNA building block functionalization; Sustainable Pharmacy

Introduction

“Old fashioned” drugs are highly active, but their lack of specificity and sensitivity needs high doses of application correlating with adverse reactions. The differentiation between tumorigenic and the surrounding healthy tissue is hardly possible. Whereas old drugs enter the cells by diffusion, the transfer of nucleic acid drugs across cell membrane is very poor and insufficient. Modern drugs and diagnostics overcome the mentioned handicaps. Therefore

a carrier system is indispensable for facilitating the transport of nucleic acid based drugs and imaging and therapy components across the cell membrane. Considerations for the improved membrane transport resulted in a series of procedures. The question respecting the low stability of nucleic acids in biological systems led to the development of numerous DNA analogues possessing higher stability.

Also important was the search for methods to
connect the carrier molecules to the therapeutic DNA derivatives or and to the intracellular contrast agents (CA) dedicated for imaging of cellular metabolic processes, combined with image-guided therapeutic approaches.

For the development of modern drugs, a very efficient ligation methodology is the Diels Alder Reaction (DAR) which traces back to 1948 and its potential and the synthesis’s mechanisms are well documented [1-3]. The DAR with inverse-electron-demand (DARinv) was first described almost 10 years later [4-7]. It is characterized by the rapid reaction rates, complete chemical reaction, lack of reverse reaction, chemical reaction at room temperature, and no need for a catalyst. Therefore the DARinv can be considered as a suitable ligation technology. Here we developed monomers based on the peptide nucleic acid’s (PNA) polyamide backbone [8], mimicking exactly the Watson-Crick hydrogen-bond formation [9-14]. The functionalization of the “PNA” like amide backbone with imaging molecules suggests a new class of efficient tools suitable for Molecular Imaging and molecular therapeutics not restricted to the classical antisense and antigenic approaches.

Here we present the synthesis of polyamide backbone pentamers and heptamers ligated with the DARinv reaction partners, fulfilling the above mentioned needs. Indeed we like to emphasize that the chemical procedures are documented [15] but in order to achieve a better understanding, the precise steps of the different chemical procedures are described particularly with full details to permit the development of modern therapeutic drugs and diagnostic molecules.

**Chemical Procedures**

1. Pentenoic acid chloride and cyclopentene carboxylic acid chloride were purchased from Sigma Aldrich, Germany. The synthesis of the Reppe Anhydride was carried out as documented by Reppe [16]. The reaction to tetracyclo-[5.4.2.7.02,6.08,11]3,5-dioxo-4-aza-9,12-tridecadiene 4-yl acetyl acid chloride is described by Wiessler [15].

2. The syntheses of the amide backbone monomers (PNA-like but without nucleobase). All synthesis steps of the tested Fmoc-protected building blocks were performed according Atherton’s and Sheppard’s [17] and Wiessler’s documentations (Figure 1) [15].

3. The synthesis of the tetrazine dicarbonic acid derivate was performed as described by [15]. The synthesis procedure of the corresponding dansyl derivate was carried out according to the following protocol (Figure 2):

![Figure 1](http://www.medscl.org)

Figure 1 illustrates the amide-based building blocks Fmoc-N-protected glycine- tert-butylester cyclopentane 1, butene 2, and Reppe anhydride 3 derivatized respectively. The synthesis of these was performed according the general procedure for the reaction of the Fmoc-glycine with acid chlorides published by Thomson [18].
4. For the syntheses of the polyamide-based pentamers I-III (Figure 3, Figure 4) and the heptamer (the ligation product 15 is shown in Figure 7) the solid phase peptide syntheses and the protection group strategies were used as introduced by Merrifield [20] and Carpino [21] considered as general procedure:

To perform the solid phase peptide synthesis (SPPS) [20] of amide modules we employed the Fmoc-strategy [21] in a fully automated peptide synthesizer A433 (Perkin Elmer). The synthesis was carried out on a 0.05 mmol Tenta Gel R Ram (Rapp Polymere) 0.19 mmol/g by substitution. As coupling agent 2-(1H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU) was used. A typical synthetic cycle consisted of a single 30 minute coupling step of 3 equivalents of monomers to the growing polyamide chain, followed by capping of the unreacted free amines with acetic anhydride. The protected polyamide resins were treated with 20% piperidine in dimethylformamide over 5 minutes and then washed thoroughly with dimethylformamide. Cleavage and deprotection of the resins were made by treatment with 90% trifluoroacetic acid and 10% triethylsilane.

5. Solid phase synthesis of the Reppe Anhydride polyamide pentamer I. To demonstrate the high efficiency of the DARinv-based “Click”-chemistry, we synthesized a pentamer which is amide-based and functionalized with the “Reppe Anhydride” 8 as shown in Figure 3: Mass spectrum: m/e 1723.3 calc. 1722.7. The corresponding ESI MS is shown in the Figure S3.

Figure 2 demonstrates the chemical reaction of the 2-cyano-nicotinic acid 4 with hydrazine to 5. After oxidation the dihydrotetrazine product bis-2,6-[5-carboxylic acid-pyrid-2-yl]-dihydropirazone 5 was transformed to the corresponding tetrazine 6, which in turn reacts with a mixture of the dansyl derivative and the N-ethyl-disoproylamine and the dansyl sulfamidoethylamine to the tetrazine product 7 linked with 5-dansyl sulfamidoethylcarboxamide-2-yl.

Figure 3 exemplifies the chemical structure of the pentamer consisting of the “Reppe Anhydride” derivative 8.
SPPS of the pentenoyl-pentamers II & the mixed Reppe Anhydride pentamers III: We also produced mixed pentamers composed of the Reppe Anhydride building block 3 (m/e 928.3 calc. 927.5) and of the pentenoic acid 2 building block 2. (Mass spectrum: 1667.5, calc. 1665.9) for chemical reaction by the solid phase peptide synthesis. (Figure 4)

9. Ligation of the pentamer I with the tetrazine-dicarboxylate 6: One µmol of the pentamer I 8 (Figure 3) and 5.5 µmol of the tetrazine 6 (intermediate as shown in Figure 2) [22] were mixed in 0.5 ml chloroform. After 10 min the red colour disappeared. After 30 min the solvent was evaporated. Mass spectrum calc. 2573.9 found m/e 1288.5 for the dication. No signal was found for the 4-fold adduct. By using 5 µmol of the tetrazine 6 the 4-fold adduct could be seen after 30 min in the mass spectrum.

10. Ligation of the pentamer I with the dansyl-tetrazine 7: One µmol (1.72 mg) pentamer I 8 (Figure 3) and 5.5 µmol (4.81mg) 7 (the reaction product is shown in Figure 2) were reacted in 0.5 ml DMSO for 12 hours. The mass spectrum showed the product at m/e 5958.9 calc. 5958.1, the trication at m/e 1986.5 and the tetracation at 1489. The dansyl-tetrazine could be seen at m/e 875.7.

11. Ligation of the Pentamer II with the tetrazine-dicarboxylate 6: The DARinv of the pentenoyl-pentamer II 2 with tetrazine-3,6-dimethylcarboxilate 2 µmol (1.86 mg) of the pentamer 8 and 10 µmol (2mg) of the tetrazine 6 in 0.5 ml chloroform were reacted for 12 hrs. The mass spectrum showed the 5-fold adducts at m/e 1779.0 calc. 1778.4, the dication at m/e 890.0. At m/e 1608.9 a weak signal appeared for the 4-fold adduct.
12. **Ligation of Pentamer II with the dansyl-tetrazine**. Two μmoles (1.86 mg) of the pentamer II (Figure 4) and 10 mmol (8.8 mg) of the dansyl-tetrazine were dissolved in 0.5 ml chloroform/DMSO and reacted for 24 hours. The mass spectrum showed m/e 5160.1 calc. 5162.9 for the 5-fold adduct, m/e 4312.9 calc 4315.6 for the 4-fold and m/e 3466.9 calc. 3469.3 for the 3-fold adduct.

13. **Ligation reaction of a polyamide heptamer III functionalized with different reactive dienophiles with two different tetrazines**. The sequence of the ligation reactions A and B are shown in Figure 7.

A. 1.66 mg (1 μmol) of the polyamide heptamer were pre-filled and reacted with 0.396 mg (2.0 μmol) of dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate 6 dissolved in 0.5 ml dichloromethane. Under stirring the reaction vessel was kept until the decolorization was complete in about 10 min.

The mass spectrum of the 2-fold adduct was at m/e 2008.7 calc. 2007.0 and the dication at m/e 1004.9. The 3-fold adduct could be seen as dication at m/e 1090.0 calc. 1089.5.

B. In a second step the above mentioned probe was reacted with 1 μmol (0.87 mg) of the dansyl-tetrazine solved in a few drops of DMSO. The reaction was completed over night. In the mass spectrum the product can be seen as dication at m/e 1428.8, corresponding to MW 2855.6; calc. 2853.2.

**Ligation Results**

Our amide building blocks, deriving from PNA devices work in a variety of ligation areas as illustrated in the following:

Using the solid phase peptide synthesis (SPPS) we could manufacture functional and modularly composed polyamides for coupling different active agents. These could be used either in parallel as imaging molecules or in combination with transporter molecules in order to reach local concentrations which were unachievable until now.

The synthesized pentamer 8 consisting of five amide-based backbone functionalized with the Reppe Anhydride derivative 3 acts as a cargo and is the reaction partner, a dienophile compound for substances harbouring diene reaction groups. The following features predispose the amide-based building block functionalized with the “Reppe Anhydride” derivative for successful use in the DAR$_{inv}$ chemistry [16].

The well controlled different reactivity of the pentenoyl group compared with the dienophile groups in the amide based monomer functionalized with the “Reppe Anhydride” allows the synthesis of polyamide oligomers consisting of two or more different dienophiles suitable for two or more independent Diels Alder Reactions with inverse-electron-demand (DAR$_{inv}$) as shown exemplarily in Figure 4.

**DAR$_{inv}$ ligation of the tetrazine derivatized polyamide pentamers**

The first highly active part of the construct, allows the ligation of e.g. carrier molecules on the desired side of the molecule. The second dienophile on the other side with lower reactivity is available for further selective functionalizations under different reaction conditions e.g. acting as coupling side for fluorescent markers.

**Ligation of the (RE-PA)$_3$ pentamer with the di-dansyl-diaryl-tetrazine**

This DAR$_{inv}$ mediated reaction describes the final product 10 of the complete ligation of the Reppe Anhydride pentamer with the dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate functionalized with two dansyl chlorides resulting in a symmetric molecule as illustrated in Figure 6.

**Ligation reaction of a polyamide heptamer functionalized with different reactive dienophiles**

A ligation of a polyamide heptamer 12 consisting of different reactive dienophiles like the Reppe Anhydride” derivative 3 and the pentenoic acid 2 is shown in Figure 1. They could also be separated by a cyclopentane building block 1, which avoids possible steric interactions restricting the ligation efficiency. The ligation starts with the chemical reaction of the Reppe Anhydride derivative monomer 3, the reaction partner with the higher reactivity. After completion, the second ligation reaction with the pentenoic acid 2 begins. The process of the chemical reaction can be monitored by the colour change and the end of the reaction is indicated by decolorization after few minutes.
Figure 5 shows the amide backbone-based pentamer Reppe Anhydride derivative functionalized after DAR$_{inv}$ mediated ligation with dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate 10. The DAR$_{inv}$ mechanism is described in detail [5].

Figure 6 shows the ligation product 11 of the polyamide pentamer 8 after the complete DAR$_{inv}$ reaction with the diene compound dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate derivatized with the fluorescent dye dansyl chloride [5-(dimethylamino)naphthalene-1-sulfonyl chloride] connected with an ethylene diamine linker.
Figure 7 shows the ligation product 12 of the polyamide heptamer molecule (in two-steps generated as described in the material and methods section): first ligation step (A) with the Reppe Anhydride derivative acting as dienophile and the dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate as diene component. The second ligation step (B) was the reaction of pentenoic acid group with the tetrazine, two fold dansyl functionalized (symmetrical molecule). Avoiding sterical interactions which can hamper the ligation processes all dienophile compounds were spatially separated with an amide-based monomer functionalized with a cyclopentane group 2.

Further examples of functionalizations

Ligation of the pentenoic acid pentamer with the dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate.

Figure 8 describes the reaction product of 13 after chemical ligation by the DARinv of the pentamer consisting of five pentenoic acid monomers 2 synthesized by SPPS as a dienophile reaction partner and with the dimethyl 1,2,4,5-tetrazine-3,6-dicarboxylate as the diene component.
Ligation of the polyamide pentenoic acid pentamer with the dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate dansyl derivatized.

Figure 9 elucidates the product of 14 after the complete DAR$_{inv}$-based ligation of the penta-amide (pentenoic acid building blocks) as a dienophile component with five diene molecules of dimethyl-1,2,4,5-tetrazine-3,6-dicarboxylate which are dansyl derivatized 7.

Discussion

Efforts were made in the development of chemoselective ligation reactions resulting in multi-faced coupling methods which are documented by Hermanson [24]. For future “Click Chemistry” applications the rapid and selective ligations must fulfill dedicated requirements e.g. to be useful in living systems, generating stable educts, intermediates, and products in order to answer questions about in vivo protein localizations and interactions as well as about cell trafficking and migration activity of stem cells for instance. The Staudinger ligation is broadly considered as an eligible candidate living up to expectations [25-27]. Here we used for such selective reactions the Diels Alder Reaction with inverse electron demand (DAR$_{inv}$) which seems to be predestinated [15, 28].

The design of different polyamides composed of the “PNA” like amide backbone whose nucleobase is substituted with components suitable for DAR$_{inv}$ ligation chemistry was established in our group and is described in the methods part.

The broad spectrum of these ligation reactions is possible by the optional exchange of building blocks which in turn can be functionalized ready for Click Chemistry according to production requirements. The new molecules were functionalized, as dienophile- or diene compounds by coupling to the glycine’s N-terminus, as originally described by Nielsen in the protocols for the synthesis of the peptide nucleic acid (PNA) backbone [29, 30].

We synthesized and investigated ligations of active components e.g. the tetracyclo[5,4.2.1.7.02,6.08,11]3,5-dioxo-4-aza-9,12-tridecadien, well known as “Reppe Anhydride”[16] which is con-
sidered as a suitable candidate for the ligation reaction with DAR_{inv}. The synthesis’s steps as well as the chemical reactions were documented in 2009 by the Pipkorn group [31].

In this paper we focused our studies at first to the synthesis of the amide pentamer functionalized with the Reppe Anhydride 8 (Figure 3). In this context it is also important to note that the used Fmoc-protected building block derivatives (Figure 1) like 3 can avoid both: I) an unnecessary molecule expansion after ligation with projected functional molecules and associated undesired reactions and II) steric effects.

Furthermore, with all these functionalized amide monomers, Fmoc-protected (Figure 1), the polyamide synthesis can be carried out by use of the established solid phase peptide synthesis (SPPS). Unnecessary coupling steps can be circumvented and the corresponding products can be achieved in satisfactorily yields and in good quality.

The Figure 1 illustrates three building blocks whereof 2 and 3 are functionalized with dienophiles featuring different chemical reactivity, 1 was functionalized with the inactive cyclopentane in order to act as a “spacer” compound avoiding spatial sterical interactions which could hamper the ligation reaction.

A descriptive example for the synthesis of polymers offering the variability for independent ligation reactions based on the DAR_{inv} with dienophile groups with different chemical reactivity is illustrated in Figure 7, which represents the reaction product after the independent ligations A and B by DAR_{inv} in 12. It shows the structural formula of the amide heptamer, functionalized with four building blocks, which possess a cyclopentane group 1, which in turn separates the two monomers featuring the Reppe Anhydride 3 and the monomer with the pentenoic acid 2 in the center of the heptamer.

The DAR_{inv} based Click chemistry’s potential is high. It allows to ligate user-defined components for imaging and/or for therapy with PNA like amide based building blocks. These examples may contribute to the establishment of a platform for expanded use in future pharmaceutical applications. In this respect positive results of toxicologic studies are indispensable to qualify such components as candidates for cell- and tissue specific therapeutic approaches. They are also ideal as diagnostic tools in the increasing fields of the patient-specific therapy and in imaging of metabolic processes at the cellular level [5, 15, 23, 32]. In summary, it is fair to say, that further efforts in the development of new functional building blocks could enhance the diversity in the ligation chemistry. They also can be conductive for both for sustainable solutions in the pharmaceutical science as discussed by Kummerer [33] and in the strongly growing field of the theranostics [34].

Conflict of Interest

The authors have declared that no conflict of interest exists.

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