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Optimized reaction conditions for amide bond formation in DNAencoded combinatorial libraries

Yizhou Li^{#1}, Elena Gabriele^{#1}, Florent Samain², Nicholas Favalli¹, Filippo Sladojevich², Jörg Scheuermann¹, and Dario Neri^{1,*}

¹Department of Chemistry and Applied Biosciences, Swiss Federal Institute of Technology (ETH Zürich), Vladimir-Prelog-Weg 3, CH-8093 Zürich (Switzerland) ²Philochem AG, Libernstrasse 3, 8112 Otelfingen (Switzerland)

[#] These authors contributed equally to this work.

Abstract

DNA-encoded combinatorial libraries are increasingly being used as tools for the discovery of small organic binding molecules to proteins of biological or pharmaceutical interest. In the majority of cases, synthetic procedures for the formation of DNA-encoded combinatorial libraries incorporate at least one step of amide bond formation between amino modified DNA and a carboxylic acid. We investigated reaction conditions and established a methodology by using 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide, 1-hydroxy-7-azabenzotriazole and N,N'-diisopropylethylamine (EDC/HOAt/DIPEA) in combination, which provided conversions greater than 75% for 423/543 (78%) of the carboxylic acids tested. These reaction conditions were efficient with a variety of primary and secondary amines, as well as with various types of aminomodified oligonucleotides. The reaction conditions, which also worked efficiently over a broad range of DNA concentrations and reaction scales, should facilitate the synthesis of novel DNA-encoded combinatorial libraries.

Keywords

DNA-encoded chemical libraries; amide bond formation; synthesis; bioconjugation

DNA-encoded combinatorial libraries (DECLs) are collections of organic compounds, individually coupled to distinct DNA fragments, serving as identification barcodes.1–4 Since DNA can be efficiently amplified by the polymerase chain reaction (PCR) and read by high-throughput DNA sequencing methods, the encoding of combinatorial libraries with DNA barcodes allows both the facile identification of specific ligands to protein targets immobilized on a solid support and the convenient handling of the libraries as mixtures of compounds. Compared to High-Throughput Screening, which relies on expensive compound

^{*}Corresponding Author: dario.neri@pharma.ethz.ch Tel.: (+41) 44-633-7401. Fax: (+41) 44-633-1358.

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collections and a sophisticated management system, DECLs provide an avenue for researchers from academia or small companies to identify hits that bind to targets of pharmaceutical interest. Numerous binders have been discovered from recent DECL selections, like ADAMTS-5 inhibitors,5 tankyrase 1 inhibitors,6 XIPA inhibitors 7 and integrin lymphocyte function-associated antigen 1 (LFA-1) antagonists,8 just to name a few. 9

Various chemical strategies have been developed for the generation of large combinatorial DECLs, including DNA-templated synthesis,10,11 DNA-encoded routing,12 hybridizationbased chemical assembling 13–17and DNA-encoded solid-phase synthesis.18 In addition, the iterative assembly of sets of building blocks in a "split-and-pool" methodology, accompanied by the stepwise addition of DNA barcodes (which unambiguously identify the chemical identity of the nascent molecules), is the most generally used strategy for DECLs construction. This strategy facilitates the preparation of DECLs containing millions or even billions of compounds, starting from just few hundred building blocks and oligonucleotides. 19,20,21

In the majority of DECLs disclosed so far, at least one synthesis step involved the formation of an amide bond.4 More than 20'000 carboxylic acids can be purchased from commercial sources at moderate costs (less than \$150/g).22 The acylation of DNA-attached amines with carboxylic acids can be performed on protected DNA structures linked to Controlled Pore Glass (CPG) supports 6 or in solution, 23 followed by deprotection and HPLC purification of the individual conjugates at early stages of library construction. Additionally, for unprotected DNA-attached amines, acylation can be performed either in solution or on "pseudo-solid phase" (i.e., on amino-modified oligonucleotide derivatives non-covalently immobilized on solid supports, facilitating the use of large molar excess of reagents and repeated washing steps for purification). 24,25 While reaction conditions with DNA on solid phase can work very efficiently, they typically consume several milligrams of each carboxylic acid for each coupling, which makes synthesis expensive and may limit the use of precious building blocks for the construction of different chemical libraries. By contrast, the acylation in solution may allow the synthesis of large DECLs using only minute amounts of building blocks.

Amide bond-forming reactions, performed in solution with unprotected oligonucleotides, have been described using EDC/*sulfo*-NHS26 or DMT-MM as coupling reagent for DECL synthesis purposes.19,20 Recently, Satz and colleagues have explored the scope and applicability of organic reactions in aqueous solution for the construction of DECLs, reporting also a sequential acylation method with DMT-MM for multistep synthesis.27 In this letter, we report an alternative a new method (featuring EDC/HOAt/DIPEA as coupling combination), which afforded >75% conversions for the majority of the tested building blocks (423 out of 543, corresponding to 78%).

Motivated by previous experience of our laboratory in the synthesis of DECLs, which involved carboxylic acids as building blocks, we characterized the reaction of eight representative compounds, leading to very different conversion rates with DMT-MM as coupling reagent [Figure 1, entry 1]. While two carboxylic acids (CA-1 and CA-2) formed

amides with 5'-aminomodified DNA (ODN-1) with excellent conversion rates (>90%; determined by UV absorbance trace at 260 nm of UPLC), the remaining six structures exhibited suboptimal conversion (ranging between 0 and 11% for 4 out of 8 compounds). A systematic investigation for alternative coupling methods is summarized in Figure 1 and different peptide coupling reagents were tested to enhance the yield of acylation. Coupling reagents like HATU [Figure 1, entries 2 and 3] and COMU [Figure 1, entry 4] consistently provided poor conversion rates for all the reactions tested. EEDQ [Figure 1, entry 5] and TBTU [Figure 1, entry 6] afforded better conversion only for some acids, while the coupling efficacy remained very low for the other substrates. Promising results were obtained by using EDC/sulfo-NHS [Figure 1, entry 7] as coupling combination, a well established and widely used method. This led to a complete conversion for 7 out of 8 compounds, while for one carboxylic acid (CA-8) no product could be observed. Further optimization revealed a novel method, featuring the combination of EDC/HOAt/DIPEA [Figure 1, entry 8], which yielded >90% conversion for all the tested compounds. For four tested acids, the coupling efficiency decreased dramatically by using only EDC/HOAt [Figure 1, entry 9], highlighting the importance of DIPEA for proper activation. Figure 2 and Supplementary Figure S3 show representative LC and MS profiles for CA-5, one carboxylic acid that did not couple at all using DMT-MM, which however provided excellent conversion (>90%) with the EDC/ HOAt/DIPEA methodology.

In order to compare the performance of the DMT-MM and EDC/HOAt/DIPEA methods over a broader range of substrates, we performed reactions using ODN-1 and a structurally diverse set of 543 carboxylic acids. Using an optimized DMT-MM protocol, 42% of the compounds exhibited a conversion lower than 50%, while 14% had a conversion between 50% and 75%, as assessed by UPLC analysis and MS confirmation [Figure 3a]. Conversely, the EDC/HOAt/DIPEA method exhibited a conversion rate of >75% for 78% of the 543 carboxylic acids. However, no clear correlation between structural features and conversion rate was found. For instance, different types of carboxylic acids (i.e., benzoic acids; primary, secondary, and tertiary carboxylic acids; aromatic or heterocyclic acids) provided similar coupling efficiencies [Figure 3b]. The experimental findings underline the importance of screening all building blocks in model reactions before DECL construction, rather than studying coupling conditions on a small set of test compounds.

In a previous report, it was described that adduct formation of EDC and G- and Tnucleobases could occur during the formation of the acylation product by using EDC as coupling reagent on "pseudo-solid phase".25 While the impact of such possible base modifications on the performance of PCR amplification is so far unexplored, it may also compromise the integrity of the DNA-barcodes.28 We systematically checked the mass spectra of all obtained conjugates from the solution phase EDC/HOAt/DIPEA method and found no detectable EDC adducts associated with the conjugates (i.e., <5%). Supplementary Figure S4 shows an exemplary MS profile for CA-9 (trans-4-hydroxycyclohexane-1carboxylic acid)-DNA conjugate, one of the carboxylic acids that exhibited significant EDC adduct formation with the "pseudo-solid phase" method. No adduct was detected with the EDC/HOAt/DIPEA methodology.

We further tested the coupling efficiency between eight representative carboxylic acids and six DNA scaffolds, containing amines with different chemical features [Figure 4]. The DNA-conjugate of glycine (SC-1) and the DNA-conjugate linked to 4- (aminomethyl)benzoic acid (SC-2) provided fair to good conversion yields for all the eight tested acids. Good conversions were also achieved with two cyclic secondary amines SC-5 and SC-6 respectively.

Using our EDC/HOAt/DIPEA coupling protocol, conversions of amide bond forming reactions were distinctly lower for sterically hindered substrates (SC-3). More steric bulk at the α -carbon further decreased the coupling efficiency extensively. For example, poor coupling results were obtained with a DNA-conjugate of 2-aminoisobutyric acid (SC-4). With the exception of CA-1, all the remaining seven carboxylic acids had conversion yields <30% under the optimized conditions.

In order to further investigate the dependence of reaction yields on the nature of amines on DNA, five additional amino-modified oligonucleotides bearing different tethers were used. Such reactions are particularly useful for the construction of encoded self-assembling chemical libraries.17 Compared to ODN-1, which featured a long C-12 tether, amino-modified oligonucleotides (i.e., ODN-2, ODN-3 and ODN-4) with various short linkers at the 5' end were examined. Good conversions were observed in 22/24 reactions and, in all cases, a conversion >60% was achieved [Figure 5]. ODN-5, carrying an aminohexyl-tether at the 3' end, was also efficiently converted into the corresponding acylated conjugates (>75%). As the chemical modification of internal sites within the DNA structure may be used for the construction of DECLs using the Yoctoliter reactor method, 29 the conversion of a primary amine on the side chain of a thymine modified at the C-6 position in ODN-6 was investigated and led to good yields for 7 out of 8 reactions, while only one carboxylic acid (CA-8) yielded a slightly lower conversion (66%).

Importantly, the EDC/HOAt/DIPEA method worked well also at different scales of aminomodified DNA (ranging between 2 and 10 nmol), which are customarily utilized for library construction [Figure 6].

The execution of synthetic steps with high yields and good purity is particularly important for the construction of DNA-encoded combinatorial libraries that require multiple steps of assembly (e.g., those based on split-and-pool methodologies). While individual DNA conjugates can be purified and analyzed after the first step of library construction, the processing of synthetic intermediates at later stages becomes more complicated. In order to minimize the presence of incomplete reaction products (e.g., residual unreacted amines) in the library, methods have been developed for the modification and removal of such truncated products.30 While these methods can be useful and may lead to improved library purity, they inevitably lead to an uneven distribution of library members, thus complicating decoding procedures. The method described in this letter should facilitate library synthesis without the need for "cap-and-catch" strategies.

In summary, we have systematically investigated different experimental conditions for amide bond formation on DNA in solution phase and established a novel efficient methodology

(EDC/HOAt/DIPEA as coupling combination). We tested 543 structurally diverse carboxylic acids as building blocks and found that the method provides good conversion yields for the majority of the compounds. The reaction works well with various types of amino-modified oligonucleotides and at different reaction scales. While the EDC/HOAt/DIPEA method worked well in most experimental conditions, additional research efforts will be needed for certain substrates (e.g., amines featuring a substantial steric hindrance), as chemical purity greatly contributes to the performance of DECL technology.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

CA	carboxylic acid
COMU	(1-cyano-2-ethoxy-2-oxoethylidenaminooxy)-dimethylamino-morpholino- carbenium hexafluorophosphate
DIPEA	<i>N</i> , <i>N</i> ['] -diisopropylethylamine
MSO	dimethyl sulfoxide
DMT-MM	4-(4, 6-dimethoxy-1, 3, 5-triazin-2-yl)-4-methylmorpholinium chloride
EDC	1-ethyl-3- (3-dimethylaminopropyl) carbodiimide
EEDQ	2-Ethoxy-1-ethoxycarbonyl-1,2-dihydroquinoline
HATU	1-[Bis(dimethylamino)methylene]-1H-1,2,3-triazolo[4,5-b]pyridinium 3- oxid hexafluorophosphate
HOAt	1-hydroxy-7-azabenzotriazole
UPLC	Ultra-performance liquid chromatography
LC-MS	liquid chromatography mass spectrometry
NMM	N-methylmorpholine
ODN	oligonucleotide
SC	scaffold conjugate
sulfo-NHS	N-hydroxysulfosuccinimide

- **TBTU** O-(Benzotriazol-1-yl)-*N*,*N*,*N*,*N*'-tetramethyluronium tetrafluoroborate
- **TEAA** triethylammonium acetate
- **TEA** triethylamine

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		DMT-MM	HATU/ DIPEA	HATU/ HOAt/ DIPEA	COMU	EEDQ	TBTU/ NMM	EDC/ sulfo-NHS	EDC/HOAt/ DIPEA	EDC/HOAt
		entry 1	entry 2	entry 3	entry 4	entry 5	entry 6	entry 7	entry 8	entry 9
но С	A-1	>90%	67%	58%	10%	>90%	>90%	>90%	>90%	>90%
	A-2	>90%	34%	22%	11%	>90%	>90%	>90%	>90%	>90%
но ССССС	A-3	69%	12%	14%	0%	85%	36%	>90%	>90%	69%
	A-4	42%	0%	0%	0%	41%	0%	>90%	>90%	0%
но ЦС С	A-5	0%	0%	0%	0%	19%	24%	>90%	>90%	16%
HO HO C	A-6	11%	53%	36%	9%	36%	>90%	>90%	>90%	>90%
$\sim \sim \sim$	A-7	5%	38%	32%	0%	8%	87%	>90%	>90%	>90%
	A-8	0%	0%	0%	0%	0%	0%	0%	>90%	25%

Figure 1.

Summary of coupling conversions by using different coupling reagents with eight representative carboxylic acids. In all cases, 72 μ L ODN-1 (MOPS buffer pH 8.0, 0.5 nmol), 45 μ L carboxylic acid (60 mM in DMSO) and coupling reagent were reacted for 16h at RT, followed by an additional coupling step with the same activated carboxylic acid for 6h at RT. Coupling reagents: entry 1: 4 μ L DMT-MM (300 mM in water); entry 2: 4 μ L HATU (300 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 3: 4 μ L HATU (300 mM in DMSO), 4 μ L HATU (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 5: 4 μ L EEDQ (300 mM in DMSO); entry 6: 4 μ L TBTU (300 mM in DMSO), 4 μ L NMM (300 mM in DMSO); entry 7: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 9: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 9: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 9: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 9: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO), 4 μ L DIPEA (300 mM in DMSO); entry 9: 4 μ L EDC (300 mM in DMSO), 4 μ L HOAt (60 mM in DMSO).

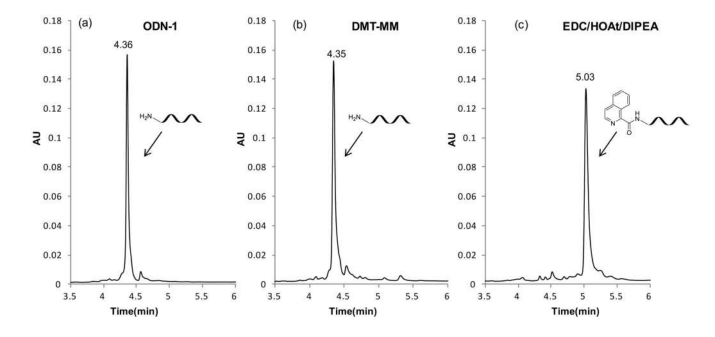


Figure 2.

UPLC profiles of (a) 5'-C12-amino oligonucleotide (ODN-1). (b) crude reaction mixture after coupling with CA-5 using DMT-MM. (c) crude reaction mixture after coupling with CA-5 using EDC/HOAt/DIPEA. Detection at 260 nm.

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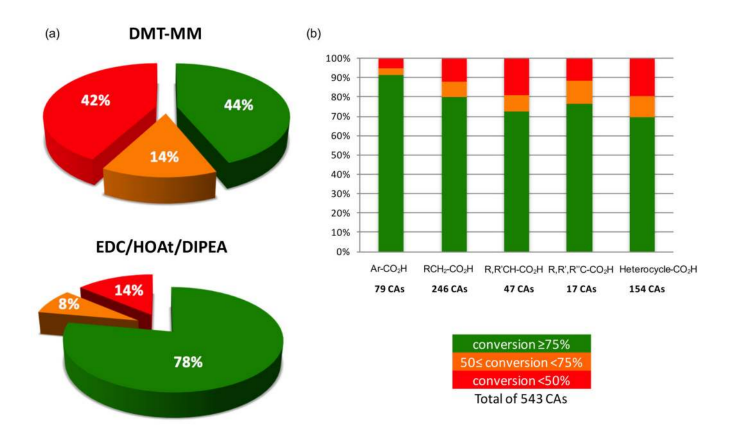


Figure 3.

(a) Pie chart analysis of the coupling yields of ODN-1 with 543 carboxylic acids using DMT-MM and EDC/HOAt/DIPEA-mediated coupling, according to the conditions described in Figure 1, entries 1 and 8. (b) Histogram analysis of the coupling yields of structurally diverse carboxylic acids to ODN-1 with EDC/HOAt/DIPEA method. Benzoic (79 acids), primary (246 acids), secondary (47 acids), and tertiary carboxylic acids (17 acids), heterocyclic acids (154 acids).

	SC-1	SC-2	SC-3	SC-4	SC-5	SC-6
CA-1	>90%	>90%	>90%	48%	>90%	>90%
CA-2	>90%	>90%	>90%	29%	>90%	>90%
CA-3	>90%	>90%	61%	0%	>90%	>90%
CA-4	>90%	89%	70%	6%	>90%	>90%
CA-5	>90%	>90%	50%	12%	75%	>90%
CA-6	>90%	>90%	>90%	17%	>90%	>90%
CA-7	>90%	87%	>90%	3%	>90%	>90%
CA-8	77%	60%	21%	5%	>90%	81%

Figure 4.

Coupling yields of six different DNA-Scaffold Conjugates with eight representative carboxylic acids, according to the condition described in Figure 1, entry 8.

	ODN-2	ODN-3	ODN-4	ODN-5	ODN-6
	00112	00110	00114	00110	
CA-1	>90%	>90%	>90%	>90%	>90%
CA-2	>90%	>90%	89%	>90%	>90%
CA-3	>90%	>90%	>90%	>90%	>90%
CA-4	>90%	>90%	>90%	79%	85%
CA-5	81%	78%	>90%	86%	76%
CA-6	>90%	>90%	>90%	>90%	>90%
CA-7	>90%	>90%	>90%	>90%	>90%
CA-8	60%	66%	80%	75%	66%

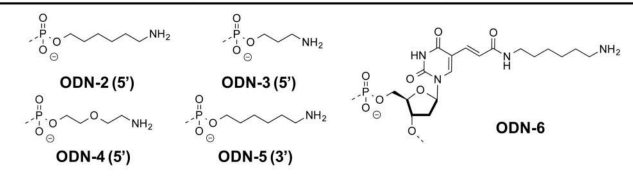


Figure 5.

Coupling yields of five different amino-modified oligonucleotides with eight representative carboxylic acids, according to the condition described in Figure 1, entry 8.

	2 nmol	10 nmol
CA-1	>90%	>90%
CA-2	>90%	>90%
CA-3	>90%	>90%
CA-4	>90%	>90%
CA-5	>90%	>90%
CA-6	>90%	>90%
CA-7	>90%	>90%
CA-8	>90%	87%

Figure 6.

Coupling yields of ODN-1 at 2 nmol and 10 nmol scale with eight representative carboxylic acids, according to the condition described in Figure 1, entry 8.