Tetrahedron Letters 58 (2017) 4391-4394

Contents lists available at ScienceDirect

Tetrahedron Letters

journal homepage: www.elsevier.com/locate/tetlet

A comparative study of amide-bond forming reagents in aqueous media – Substrate scope and reagent compatibility



Matthew Badland ^a, Robert Crook ^a, Bastien Delayre ^b, Steven J. Fussell ^{a,*}, Iain Gladwell ^a, Michael Hawksworth ^a, Roger M. Howard ^c, Robert Walton ^a, Gerald A. Weisenburger ^c

^a Pfizer Worldwide Research and Development, Discovery Park, Sandwich, Kent CT13 9NJ, United Kingdom
 ^b CPE Lyon, 43 Bd du 11 Novembre 1918, F-69100 Villeurbanne, France
 ^c Pfizer Worldwide Research and Development, Eastern Point Road, Groton, CT 06340, United States

ARTICLE INFO

Article history: Received 26 August 2017 Revised 3 October 2017 Accepted 5 October 2017 Available online 7 October 2017

Keywords: Amide-bond forming Condensation in water Coupling reagents

ABSTRACT

A survey of amidation reagents demonstrating DIC-HOPO, DMT-MM, COMU-collidine, TPTU-NMI, EEDQ, CDI and EDC-Oxyma to be effective for the coupling of carboxylic acids with amines in the presence of water and the absence of problematic dipolar aprotic solvents is reported. DMT-MM was shown to provide the best yields for the coupling of a secondary amine, TPTU-NMI and COMU-collidine for aniline, whilst the combination of DIC with HOPO afforded the broadest substrate scope and the highest yields for a sterically demanding carboxylic acid.

© 2017 Elsevier Ltd. All rights reserved.

The amide functionality is an important component of many drug molecules. A recent survey showed that amidation featured in approximately 50% of all Journal of Medicinal Chemistry manuscripts examined, making it the most frequently used synthetic transformation, and importantly with 30% as the final bond forming step.¹ Amide bonds are typically synthesised by reactions of carboxylic acids and amines with the loss of water, in most cases aided by a coupling reagent. A plethora of coupling reagents are commercially available and have been widely used in large-scale manufacture of drug candidates.^{2,3} Regrettably, undesirable dipolar aprotic solvents (particularly the reprotoxic solvents dimethylacetamide (DMAC), dimethylformamide (DMF) and N-methyl pyrrolidone (NMP) are commonly used^{2b,3} due to the poor organic solubility of carboxylic acids, carboxylate salts (including amine/carboxylic acid salt pairs) and zwitterionic substrates. A number of these solvents are deemed substances of high concern and are subject to considerable attention under European REACh regulation.⁴ Given the importance of amidation reactions, REAChunencumbered solvent systems would be highly desirable in designing new scaleable chemical processes.

It is commonly known that carboxylic acids, carboxylate salts and α -amino acids can be readily solubilized in aqueous solvent systems thus providing a potential alternative to dipolar aprotic solvents. However, for the amide coupling to succeed the rate of

* Corresponding author. *E-mail address:* Steven.Fussell@pfizer.com (S.J. Fussell). aminolysis must be significantly greater than the rate of hydrolysis of the activated carboxylic acid intermediate(s) and the coupling reagent itself. A variety of reagents have been reported to provide such conditions- most notably EDC,⁵ DPTF,⁶ DMT-MM,⁷ CDI,⁸ COMU⁹ and *N*-carboxyanhydrides (NCAs).¹⁰ Herein we describe our efforts to investigate and understand the performance of new and existing water-compatible amide coupling systems.

The study was initiated by screening 48 different coupling conditions for the amidation of benzoic acid with benzylamine in the presence of water (Fig. 1). The reaction were initially carried out using NMP as an organic co-solvent to mitigate any solubility issues and was executed by simultaneous addition of the carboxylic acid and amine to the amidation reagent in solution. A range of coupling reagents were shown to afford moderate to high *in situ* yields. Lead reagents selected for further study included a variety of carbodiimides (Table 1, entries 2–6, 8, 9, 11), triazines (Entries 14, 15, 20–22), quinoline based reagents (Entries 23, 24), COMU (Entry 25), TPTU (Entry 41), pivalic anhydride (Entry 42) and CDI (Entry 45).

Using the lead reagents described above, each system was optimized for solvent, additive, order of addition and reaction time in the model reaction (Fig. 1). Problematic dipolar aprotic co-solvents were omitted at this stage in favor of non-reprotoxic alternatives. *N*,*N*-Diisopropylcarbodiimide (DIC) in combination with HOPO as an additive, was shown to be of particular interest affording the highest *in situ* yields of product 3 (Table 2, entry 1). An EDC-Oxyma cocktail^{5b} also gave promising results, as did DMT-MM·BF₄ which





etrahedro



Fig. 1. Model reaction for the screening experiments.

Table 2

Optimized yields for the coupling of benzoic acid with benzylamine in MeCN/Water. See ESI for conditions.

Entry	Reagent	Additive	Yield (%) ^a
1	DIC	НОРО	93
2	DIC	Oxyma	80
3	DIC	HOBt·H ₂ O	81
4	EDC	Oxyma	82
5	DMT-MM·BF ₄		90
6	DMT-MM·Cl		58
7	COMU	Collidine	87
8	TPTU	NMI	90
9	EEDQ		63 ^b
10	IIDQ		79 ^b
11	Pivalic anhydride		58 ^c
12	CDI		85 ^c

^a *In situ* yield by comparison to 1,3-benzodoxole as an internal standard.

^b Addition of 4M HCl (0.1 eq.) in dioxane, 72 h reaction time.

^c Sequential activation of the carboxylic acid in acetonitrile followed by addition of the amine coupling partner as an aqueous solution.

significantly outperformed the corresponding chloro analogue (Entries 4–6). Improvement of the *in situ* yields for the TPTU and COMU systems required the addition of an organic base. *N*-Methylimidazole (NMI) in combination with TPTU (Entry 8) and collidine with COMU (Entry 7) proved to be the two best combinations of those tested, mirroring the system reported by Lipshutz and co-workers who employed a surfactant with a COMU and collidine reagent cocktail.⁹ In contrast, optimization of the quinoline-based coupling reagents indicated the requirement for sub-stoichiometric amounts of hydrochloric acid and longer reaction times (Entries 9 and 10). In the case of CDI and pivalic anhydride, a yield enhancement was achieved by sequential activation of benzoic

acid in acetonitrile followed by the addition of benzylamine as an aqueous solution (Entries 11 and 12). Additional co-solvent screening demonstrated that NMP could be readily replaced by non-reprotoxic water-miscible alternatives such as acetonitrile and tetrahydrofuran (see ESI, Section 3.1).

Following these initial studies, the substrate scope of each of the preferred amidation conditions was investigated (Table 3). The results, as expected, confirmed that the performance of each coupling reagent is substrate-dependent, underlining the importance of reagent screening when developing an amide bond forming process. COMU-collidine demonstrated one of the broadest substrate scopes, affording moderate to high yields for the coupling of both benzoic acid and 3-phenylpropanoic acid with primary and secondary amines including aniline (Table 3, entries 1-4 and 9-12). Similarly, TPTU-NMI showed broad scope and coupled aniline effectively. However, both COMU-collidine and TPTU-NMI were unable to couple the sterically hindered 2.6-dimethylbenzoic acid resulting in little or no product being formed with either amine partner tested (Entries 13 and 14). DMT-MM performed well for the coupling of dibenzylamine, generally outperforming all other coupling reagents for this amine (Entries 4, 8 and 12), yet it did not readily accept aniline or 2,6-dimethylbenzoic acid coupling partners. EEDQ and EDC-Oxyma achieved modest yields with no clear trends for substrate scope shown. The traditional reagent. CDI, was effective for the coupling of aliphatic primary amines with sterically unhindered acids (Entries 2, 3, 6, 7, 10 and 11) but only moderately so for aniline and poorly so for dibenzylamine (Entries 1, 4, 5, 8, 9 and 12). In contrast to all other reagents, the combination of DIC-HOPO was shown to perform well with all carboxylic acid and amine partners tested. Pleasingly, under forcing conditions (70 °C 2 d), DIC-HOPO even accepted the sterically demanding 2,6-dimethylbenzoic acid with both benzylamine and pyridin-2-ylmethanamine (Entries 13 and 14). However, dimethylbenzoic acid was not as well coupled to aniline and dibenzylamine, giving negligible conversions at 20 °C and thus was not optimized (results not presented).

Reaction profiling of the amidation of 2,6-dimethylbenzoic acid with benzylamine showed that the addition of HOPO with DIC

Table 1

Coupling reagent (1.0 eq.) screen for the amidation of benzoic acid with benzylamine (1.0 eq.) in NMP (21 mL/g)/water (9 mL/g) at 20 °C.

Entry	Туре	Reagent	Yield (%) ^a	Entry	Туре	Reagent	Yield (%) ^a
1	Carbodiimide	DCC	13	25	Uronium/Aminium	COMU	40
2		DCC-HOBt·H ₂ O	65	26	,	HOTU	32
3		DCC-Oxyma	63	27		HATU	52
4		DIC	65	28		HBTU	52
5		DIC-HOBt·H ₂ O	86	29		HCTU	50
6		DIC-Oxyma	81	30		HATU-HOBt·H ₂ O	49
7		EDC	4	31		HBTU-HOBt·H ₂ O	50
8		EDC-HOBt·H ₂ O	82	32		HCTU-HOBt·H ₂ O	46
9		EDC-Oxyma	83	33		HBTU-Oxyma	46
10		EDC methiodide	3	34		TPTU	49
11		EDC methiodide-HOBt·H ₂ O	79	35		HSTU	11
12		CMC	1	36		TSTU	12
13		EDC-HCl	42	37		TOTU	35
14	Triazine	DMT-MM·Cl	59	38	Phosphonium	РуАОР	32
15		DMT-MM·BF ₄	94	39		PyBrOP	12
16		Cyanuric chloride	15	40	Imidazolium	CIP	23
17		Cyanuric chloride-QD ^b	30	41		DMC	20
18		DCMT	13	42	Miscellaneous	Pivalic anhydride	59
19		DCMT-QD ^b	41	43		Benzyl chloroformate	28
20		CDMT	62	44		TFFH	20
21		CDMT-QD ^b	83	45		CDI	36
22		CDMT-DABCO	68	46		DPP	0
23	Quinoline	EEDQ	54	47		DTPC	2
24		IIDQ	42	48		TODT	25

^a In-situ yield by HPLC analysis with 1,3-benzodoxole as an internal standard.

^b Quinuclidine abbreviated to QD.

Table 3
Substrate scope for water-compatible coupling reagents. MeCN/water (1:1), coupling reagent (1.05 eq.), additive (1.0 eq.) at 20 °C.

Entry	Carboxylic Acid	Amine	DMT- MM·BF4 ^a	TPTU- NMI	COMU- collidine	Pivalic anhydride ^a	CDI ^a	EEDQ	DIC- HOPO	EDC- oxyma
			Yield (%) ^b							
1	ОН	NH ₂	15	71	79	19	14	39	69	63
2	ОН	NH ₂	82	85	80	10	85	65	89	66
3	ОН	NH2	71	76	71	7	78	57	79	56
4	ОН	N N N	85	28	69	26	1	28	68	65
5	ОН	NH ₂	57	84	81	2	71	76	78	56
6	ОН	NH ₂	62	69	47	3	74	57	74	52
7	ОН	N NH ₂	46	63	21	37	55	16	72	56
8	ОН	N H	89	76	86	5	25	76	71	70
9	ОН	NH ₂	37	85	89	41	53	74	83	54
10	ОН	NH ₂	76	88	87	39	86	73	87	60
11	ОН	NNH ₂	73	74	91	41	86	61	81	71
12	ОН	N N N	89	61	75	53	5	14	98	67
13 ^c	ОН	NH ₂	3	0	5	0	0	0	66 (60) ^d	0
14 ^c	ОН	N NH ₂	0	1	0	1	0	1	64	0

^a Sequential activation of the carboxylic acid in MeCN for DMT-MM, CDI, and Pivalic anhydride.

^b In situ yield by comparison to 1,3-benzodioxole as an internal standard.

^c Reactions with 2,6-dimethylbenzoic acid were conducted at 70 °C, 48 h.

^d Isolated yield.

completely suppressed formation of the unwanted *N*-acylurea **6**, allowing rapid conversion to the activated HOPO intermediate **7** *via* the diisopropylcarbamimidic anhydride **5** (Fig. 2).¹⁰ The formation of intermediate **7** at 20 °C in acetonitrile was complete within 5 mins and was stable to the addition of water and heating to 70 °C. Subsequent addition of benzylamine afforded rapid conversion to the desired product in moderate *in situ* yield with significant levels of residual activated HOPO intermediate **7** remaining. Continued heating of the reaction mixture at 70 °C for 1 h confirmed

that intermediate **7** was stable to the aqueous reaction medium with no regeneration of the 2,6-dimethylbenzoic acid.

To further compare the performance of the leading coupling reagents, the relative aqueous stabilities of their corresponding activated acids were investigated. Both benzoic and hydrocinnamic acid were selected as test substrates and were activated with DIC-HOPO, DIC, COMU (without additive), TPTU (without additive), CDI, DMT-MM·BF₄ and EEDQ in MeCN. Water was charged to each activated acid solution and their retained activity measured by



Fig. 2. Reaction pathway for the amidation using DIC-HOPO.

Table 4

Stability of activated benzoic and hydrocinnammic acid in aqueous media (held at 30 $^\circ$ C for 15 min in MeCN/water).

Entry	Activating agent	Retained activing intermediate	Retained activity of activated intermediate (%) ^a		
		PhCOOH	Ph(CH ₂) ₂ COOH		
1	DIC-HOPO	99	94		
2	DIC	89	83		
3	COMU	16	8		
4	TPTU	10	23		
5	CDI	85	99		
6	DMT-MM·BF ₄	49	67		
7	EEDQ	88	95		

^a The reaction mixture was quenched into *n*-butylamine and conversion to the corresponding amide was calculated.

quenching the reaction with *n*-butylamine after 15 min (Table 4). When the DIC-HOPO-activated intermediates were studied as such we were pleased to discover that only 1% of the original activity was lost by hydrolysis back to benzoic acid and only 6% lost by hydrolysis to hydrocinnamic acid (Entry 1). DIC (without additive), CDI and EEDQ activated intermediates also retained the majority of their original activity under the aqueous conditions tested (Entries 2, 5 and 7).

In conclusion, a variety of amidation reagents have been shown to promote the coupling of carboxylic acids with amines in aqueous reaction media, thus avoiding problematic dipolar aprotic solvents. The combination of DIC-HOPO provided the most impressive substrate scope and activated intermediate stability. It was also found that DMT-MM was preferred for the conversion of dibenzylamine (the only secondary amine tested) and the uronium based TPTU-NMI and COMU-collidine reagent cocktails provided the highest conversions for aniline. It is hoped that these results will encourage greater utilization of aqueous media in amide coupling and that the reliance on problematic dipolar aprotic solvents may be overcome in the pursuit of sustainable and greener chemistry.

Acknowledgments

The authors would like to thank David Daniels and Olivier Dirat for their helpful suggestions and David Mason for his analytical support.

A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.tetlet.2017.10.014.

References

- 1. Brown DG, Boström J. J Med Chem. 2016;59:4443–4458.
- 2. (a) Dunetz JR, Magano J, Weisenburger GA. Org Process Res Dev. 2016;20:140–177;
 - (b) Montalbetti CAGN, Falque V. Tetrahedron. 2005;61:10827-10852.
- (a) MacMillan DS, Murray J, Sneddon HF, Jamieson C, Watson AJB. Green Chem. 2013;15:596–600;
- (b) El-Faham A, Albericio F. Chem Rev. 2011;111:6557-6602;
- (c) Sperry JB, Farr RM, Levent M, et al. Org Process Res Dev. 2012;16:1854–1860;
 (d) Ormerod D, Willemsens B, Mermans R, et al. Org Process Res Dev. 2005;9:499–507;
- (e) Shendage DM, Fröhlich R, Haufe G. Org Lett. 2004;6:3675-3678.
- European Chemicals Agency (ECHA), Candidate list of substances of very high concern for authorisation, available online at http://echa.europa.eu/candidatelist-table. 1-Methyl-2-pyrrolidone (NMP), Toxic for reproduction (Article 57c) sampled on the 30th June 2017 (a) Konwar M, Ali AA, Sarma D. Tetrahedron Lett. 2016;57:2283–2285;
 - (b) Wang Q, Wang Y, Kurosu M. Org Lett. 2012;14:3372–3375;
 - (c) Galanis AS, Albericio F, Grotli M. Org Lett. 2009;11:4488-4491;
 - (d) Hojo K, Ichikawa H, Fukumori Y, Kawasaki K. Int J Pept Res Ther. 2008;14:373–380;
 - (e) Shi Y-J, Cameron M, Dolling UH, et al. Synlett. 2003;5:647-650;
 - (f) Nakajima N, Ikada Y. Bioconjug Chem. 1995;6:123-130;
 - (g) Ho GJ, Emerson KM, Mathre DJ, Shuman RF, Grabowski EJJ. J Org Chem. 1995;60:3569-3570;
- (h) Sheehan JC, Hlavka J. J Org Chem. 1956;21:439-441.
- 5. Murakami M, Hayashi M, Tamura N, Hoshino Y, Ito Y. Tetrahedron Lett. 1996;37:7541-7544.
- (a) Mizuhara T, Hioki K, Yamada M, Sasaki H, Morisaki D, Kunishima M. *Chem* Lett. 2008;37:1190–1191;
 (b) Hojo K, Maeda M, Tanakamaru N, Mochida K, Kawasaki K. Protein Pept Lett.
- (b) Hojo K, Maeda M, Falakamaru N, Mochida K, Kawasaki K. Protein Pept Lett.
 2006;13:189–192;
 (c) Kunishima M, Kawachi C, Hioki K, Terao K, Tani S, Tetrahedron.
- 2001;57:1551–1558.
- 7. (a) Verma SK, Ghorpade R, Pratap A, Kaushik MP. Tetrahedron Lett. 2012;53:2373–2376;
 (b) Padiya KJ, Gavade S, Kardile B, et al. Org Lett. 2012;14:2814–2817.

(D) Padiya KJ, Gavade S, Kardile B, et al. Org Lett. 2012;14:2814–2817.

- Gabriel CM, Keener M, Gallou F, Lipshutz BH. Org Lett. 2015;17:3968–3971.
 De Marco R, Tolomelli A, Greco A, Gentilucci L. ACS Sustainable Che. Eng. 2013:1:566–569.
- (a) Iwasawa T, Wash P, Gibson Ch, Rebek J. Tetrahedron. 2007;63:6506–6511;
 (b) Mikolajczyk M, Kielbasinski P. Tetrahedron. 1981;37:233–284.