RSC Advances



PAPER

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Cite this: RSC Adv., 2023, 13, 888

New boro amino amide organocatalysts for asymmetric cross aldol reaction of ketones with carbonyl compounds†

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Distinct types of new boron fused primary amino amide organocatalysts were designed and synthesized from commercially available amino acids. Their catalytic activities were investigated in asymmetric crossed aldol reaction of ketones with aromatic aldehydes to afford the corresponding chiral *anti*-aldol adducts with good chemical yields, moderate diastereoselectivity and good to excellent enantioselectivities (up to 94% yields, up to 90:10 dr, up to 94% ee).

Received 5th October 2022 Accepted 20th December 2022

DOI: 10.1039/d2ra06272k

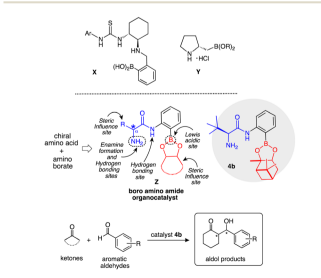
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1. Introduction

The rapidly growing interest in the area of asymmetric organocatalysis is not only due to the versatile character of small organic molecules which function as efficient and selective catalysts but is also attributed to their important role in the construction of complex and enantiopure molecular skeletons.1 Several contributions to the study of asymmetric aldol reactions were reported earlier² and with the development of the proline catalysed direct intermolecular asymmetric aldol reaction by List, Lerner and Barbas III,3 the synthesis and application of organocatalysts based on enamine-iminium ion activation and hydrogen bonding activation has received further considerable attention.4 Most organocatalysts currently used are functionally synergic systems which have two or more distinct functionalities within the same molecule. Among the various polyfunctional organocatalysts available, one functional group works as enamine-iminium formation or Brønsted basic sites and the other works as a Lewis acidic site.5 Our group is also exploring new polyfunctional organocatalysts based on primary amino alcohols and amino amides, which are easily derived from commercially available amino acids, in numerous asymmetric reactions.6 However, only a few examples have been reported on the polyfunctional organocatalysts with boron functional

Based on the above background, we designed a boron derived amino amide organocatalyst \mathbf{Z} , obtained by facile synthetic transformation. Catalyst \mathbf{Z} contains an amino group acting as an enamine formation site, amino amide group for hydrogen bonding site, boron atom acting as a Lewis acidic site, and also pinene and the β -position substituent act as steric influence sites in a single molecule (Scheme 1). The cross aldol reaction of ketones with aldehydes was chosen to explore the activity of \mathbf{Z} as an organocatalyst. The aldol reaction β is one of the most versatile and popular method for the formation of C–C bonds β 0 in modern organic synthesis.

Herein, we describe new boro amino amide Z as an organocatalyst, which showed an efficient catalytic activity in



Scheme 1 Asymmetric cross aldol reaction of ketones with carbonyl compounds.

groups as Lewis acidic sites, such as thiourea-boronic acid **X** by Takemoto⁷ and proline boronate **Y** by Whiting,⁸ so far.

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 $[\]dagger$ Electronic supplementary information (ESI) available: Experimental details and HPLC data. See DOI: https://doi.org/10.1039/d2ra06272k

crossed aldol reaction of ketones with aromatic aldehydes to afford aldol products. Especially, catalyst **4b** showed satisfactory catalytic activity in this reaction to afford the corresponding aldol products (up to 94%, up to *anti*: *syn*/90:10, up to 94% ee) in eco-friendly sea-H₂O-tap H₂O solvent condition.

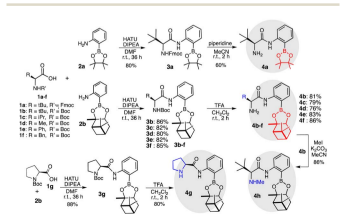
2. Results and discussion

2.1. Catalyst design and synthesis

Initially, we synthesized catalysts **4a–f** having primary amino group by the coupling of protected amino acids **1a–f** with amines **2a** and **2b** having boron substituents, respectively, followed by the deprotections of protected boro amino amides **3a** and **3b–f**, respectively, in good yields (Scheme 2). Moreover, secondary amino catalyst **4g** having proline ring was synthesized by condensation of **1g** with **2b**, followed by the deprotection of **3g**. Catalyst **4h** having secondary amino group was derived from **4b** by methylation with MeI and K₂CO₃.

2.2. Catalytic efficiency

To find out the best catalyst, our preliminary investigations were focused on screening of the synthesized catalysts 4a-h in the crossed aldol reaction of cyclohexanone 5a as an aldol donor with p-nitro benzaldehyde 6a as an aldol acceptor (Table 1). First, the catalytic activity of boro amino amide 4a with primary amino group and pinacolyl boron substituent was examined under THF at room temperature, but the formation of the corresponding aldol product 7a or 8 was not observed (entry 1). Next, this reaction using benzoic acid as a co-catalyst was examined (entries 2-9). The reaction was carried out with 10 mol% of 4a and 20 mol% of benzoic acid in THF at room temperature. Initially, the functionality of catalyst 4a was examined (entry 2). However, this catalyst 4a did not work enough and the corresponding aldol product 7 was obtained in low chemical yield as racemate. Next, catalysts 4b-f with primary amino group, several substituents at α-position and bulky pinenyl boron group on amide amino group were investigated (entries 2-7). As a result, all the catalysts showed catalytic activity. Especially, the use of catalyst 4b with tert-butyl group afforded anti-aldol product 7a as a main product with



Scheme 2 Preparations of catalysts 4a-h.

Table 1 Asymmetric aldol reaction of 5a and 6a with organocatalysts 4a-h

Entry	Cat.	Co-cat.	Yield ^a (%)	dr ^b anti:syn	ee ^c (%)
1	4a	_	nr	_	
2	4a	Benzoic acid	15	58:42	racemic
3	4b	Benzoic acid	24	60:40	66
4	4c	Benzoic acid	12	54:46	33
5	40 4d	Benzoic acid	22	55:45	40
6	4u 4e	Benzoic acid	20	58:42	27
7	4f	Benzoic acid	13	33:67	20
8	4g	Benzoic acid	28	60:40	56
9	4h	Benzoic acid	nr	_	_

^a Isolated yields (a mixture of diastereomer). ^b The dr vales were determined by ¹H NMR. ^c The ee values were determined by HPLC analysis (Daicel Chiralpak AD-H column).

moderate diastereo and enantioselectivity (anti:syn/60:40, 66% ee), although chemical yield was low (24%) (entry 3). We also screened the activity of secondary amino catalyst 4g having pyrrolidine ring that is valid for forming enamine with ketone 5a (entry 8). However, the sufficient improvement of both chemical yield and stereoselectivities (28%, anti:syn/60:40, 56% ee) in the reaction using 4g was not observed in comparison with the result of catalyst 4b having primary amino group. Based on the above results, the functionality of catalyst 4h which is obtained by a change of primary amino group on 4a to secondary amino group was also examined under similar reaction condition (entry 9). However, contrary to expectation, catalyst 4h did not work in this reaction. In addition, the aldol reaction was also carried out in the similar conditions as reported by Whiting et al., using the HCl salt of our best catalyst 4b, but 4b HCl salt catalyst only showed less catalytic activity and the aldol product 7a was observed as racemate (Scheme 3). From all of the above results, it was observed that catalyst 4b with primary amino group and both tert-butyl group at α -position and bulky pinenyl boron group on amide amino group as a steric influence sites performed better comparatively with other catalysts 4a,c-h. The absolute configuration and anti/syn diastereoselectivity of 7a and 8 were identified based on comparison with literature data.11

2.3. Improving catalytic efficiency

To improve the catalytic efficiency, the reaction conditions such as solvent, co-catalysts, and reaction times were examined by

Scheme 3 Aldol reaction under Whiting et al. condition.

using superior catalyst 4b (Table 2). First, solvent effect was examined with THF, i-PrOH, MeOH and H₂O as polar protic solvent, DMF as a polar aprotic solvent, dioxane as ethereal solvent (entries 1-6), neat condition (entry 7) and CH₂Cl₂ as halogenated solvent (entry 8) in the presence of catalyst 4b (10 mol%) and benzoic acid as a co-catalyst (20 mol%) at room temperature for 96 h. As a result, only tap H₂O had a quite influence on the enantioselectivity of the reaction to afford antialdol product 7a with moderate chemical yield and diastereoselectivity (50%, anti:syn/64:36, 85% ee) (entry 4). It was observed that based on the previous reports¹² of aldol reaction in water, which suggest that water molecule facilitates the ease of the condensation process and enhances the reactivity of the reaction, with no less influence on the transition state, especially in the asymmetric aldol reaction. Next, a series of cocatalysts were examined using superior tap H2O as solvent (entries 9-12). With all the co-catalysts, the results were found to be inferior to benzoic acid in terms of chemical yield and stereoselectivity suggesting benzoic acid as superior co-catalyst. Next, the reaction was carried out in different types of H₂O (Table 3). Initially, sea H₂O was used in this reaction and the product 7a was obtained with moderate enantioselectivity (60% ee), but chemical yield was increased to 86% with moderate diastereoselectivity (anti:syn/68:32) (entry 1). Using distilled H₂O, chemical yield was low (48%), but the enantioselectivity was relatively good (77% ee) with moderate diastereoselectivity (anti: syn/67: 33) (entry 2). The use of brine afforded 7a with moderate chemical yield and stereoselectivities (50%, anti: syn/ 62:38, 67% ee) (entry 3). From those results, it was indicated that sea H₂O is effective to increase the chemical yield of the aldol product 7a. This might be due to the high density, higher electrical conductivity, slight basic nature of sea water in

not clear. We then changed the time duration of the reaction using sea water for 72 h and 120 h. As a result, 72 h of reaction time was better for obtaining 7a in good chemical yield, moderate diastereoselectivity and enantioselectivity (80%, anti: syn/68:32, 75% ee) (entry 4). On further increase of reaction time to 120 h resulted in the aldol product in low enantioselectivity (90%, anti: syn/66:34, 11% ee) (entry 5).

Good yield was observed with sea water and relatively good enantioselectivity was observed with tap water. Therefore, we attempted the aldol reaction in a mixture of sea H₂O and tap $H_2O(1:1)$ while varying the co-catalysts and catalyst mol ratio. Initially, the reaction using mixed sea H2O-tap H2O with 10 mol% of catalyst 4b and 20 mol% of benzoic acid as a cocatalyst was investigated (82%, anti: syn/68:32, 94% ee) (entry 6). To our surprise, the aldol product 7a was obtained in good enantioselectivity (94% ee), good chemical yield and moderate diastereoselectivity (82%, anti: syn/68:32). Further, on changing the catalyst and co-catalyst mol ratios (entries 7-10) with the solvent system as sea H2O-tap H2O, the obtained results were found to be inferior to the result of entry 6. Based on the above results, the catalyst 4b (10 mol%) and benzoic acid (20 mol%) as a co-catalyst in the mixed solvent of sea H₂O-tap H₂O at 72 h was confirmed as the optimum condition.

2.4. Substrate scope

After the optimization of reaction conditions, we examined the generality of the superior catalyst **4b** in the aldol reactions of various ketones **5a-h** with aldehydes **6b-k** (Scheme 4). The reactions were carried out in the presence of catalyst **4b** (10 mol%) and benzoic acid as a co-catalyst (20 mol%) under the best reaction condition (sea H_2O -tap H_2O , 72 h, 25 °C).

Initially, substituent influences on the aromatic ring of aldehydes **6b**-**j** were examined. The reactions of **5a** with nitro aromatic aldehydes **[6b** (*m*-NO₂), **6c** (*o*-NO₂)] gave the

Table 2 Optimisation of reaction conditions using catalyst 4b

combination with mineral contents were assumed to accelerate

the reaction yields of the aldol reaction, although the reason is

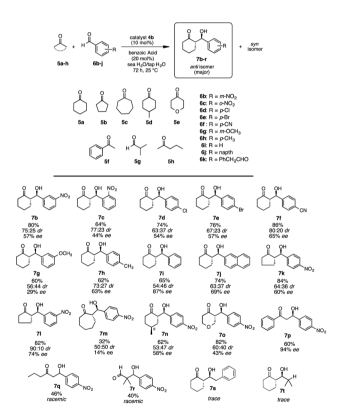
Entry	Solvent	Co-cat.	$Yield^{a}$ (%)	dr^b anti: syn	ee ^c (%)
1	THF	Benzoic acid	24	60:40	66
2	i-PrOH	Benzoic acid	20	63:37	53
3	МеОН	Benzoic acid	25	52:48	67
4	Tap H ₂ O	Benzoic acid	50	64:36	85
5	DMF	Benzoic acid	10	55:45	51
6	1,4-Dioxane	Benzoic acid	trace	_	_
7	Neat	Benzoic acid	51	54:46	59
8	CH_2Cl_2	Benzoic acid	20	61:39	60
9	Tap H ₂ O	4-NO₂Benzoic acid	45	58:42	40
10	Tap H ₂ O	TFA	50	66:33	60
11	Tap H ₂ O	Acetic acid	41	60:40	72
12	Tap H ₂ O	2-F-Benzoic acid	40	63:37	54

^a Isolated yields (a mixture of diastereomer). ^b The dr ratios were determined by ¹H NMR. ^c The ee values were determined by HPLC analysis (Daicel Chiralpak AD-H column).

Table 3 Optimisation of reaction condition using H₂O solvent using 4b

Entry	Solvent	Time (h)	Cat. (mol%)	Co-cat. (mol%)	Yield ^a (%)	$\mathrm{dr}^b \ anti: syn$	ee ^c (%)
1	Sea H ₂ O ^d	96	10	20	86	68:32	60
2	Distilled H ₂ O	96	10	20	48	67:33	77
3	Brine H ₂ O	96	10	20	50	62:38	67
4	Sea H ₂ O	72	10	20	80	68:32	75
5	Sea H ₂ O	120	10	20	90	66:34	11
6	Sea H ₂ O/tap H ₂ O	72	10	20	82	68:32	94
7	Sea H ₂ O/tap H ₂ O	72	20	20	94	43:57	64
8	Sea H ₂ O/tap H ₂ O	72	20	10	90	68:32	72
9	Sea H ₂ O/tap H ₂ O	72	10	10	44	67:33	71
10	Sea H ₂ O/tap H ₂ O	72	5	10	30	69:31	83

a Isolated yields (a mixture of diastereomer). The dr ratios were determined by HNMR. The ee values were determined by HPLC analysis (Daicel Chiralpak AD-H column). d Sea H₂O source: Itanki beach, Pacific sea, Higashi Muroran, Hokkaido, Japan. Tap H₂O: obtained from the laboratory tap H₂O.



Scheme 4 The aldol reactions of 5a-h with different aromatic aldehydes 6b-k.

corresponding aldol products 7b,c respectively, with moderate to good chemical yield and diastereoselectivities, but with moderate enantioselectivities (7b: 80%, anti: syn/75: 25 dr, anti: 57% ee, syn: 58% ee; 7c: 64%, anti: syn/77: 23 dr, anti: 44% ee, syn: 25% ee). Whereas the reaction of 5a with halogenated aromatic aldehydes [6d (p-Cl), 6e (p-Br)] gave the corresponding aldol products 7d,e respectively, with good chemical yields,

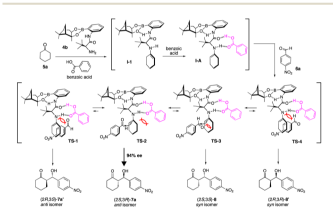
diastereoselectivities and moderate enantioselectivities (7d: 74%, anti: syn/63:37 dr, anti: 54% ee, syn: 10% ee; 7e: 76%, anti: syn/67: 23 dr, anti: 57% ee, syn: 16% ee). Furthermore, the reactions of 5a with aldehyde 6f (p-CN), having electron withdrawing group afforded the aldol product 7f in good chemical yield and moderate to good diastereoselectivities and moderate enantioselectivity (7f: 86%, anti: syn/80: 20 dr, anti: 65% ee, syn: 60% ee). The influence of electron donating group on aldehydes were also examined by the reactions of 5a with 6g (m-methoxy) and with 6h (p-methyl) to afford the corresponding anti-aldol products 7g and 7h in moderate chemical yields, moderate to good diastereoselectivities and low to moderate enantioselectivities (7g: 60%, anti: syn/56: 44 dr, anti: 29% ee, syn: 34% ee; 7h: 62%, anti: syn/73:27, anti: 63% ee, syn: 12% ee), respectively. When benzaldehyde 6i and bulkier napthaldehyde 6j were used, the chemical yields, diastereoselectivities and enantioselectivities were moderate to good (7i: 55%, anti: syn/ 54: 46 dr, anti: 87% ee, syn: 12% ee; 7j: 74%, anti: syn/63: 37 dr, anti: 69% ee, syn: 40% ee). Furthermore, the reactions of 5 membered cyclopentanone 5b with 6a,b and 7 membered cycloheptanone 5c with 6a, respectively, was carried out in the presence of catalyst 4b (10 mol%) and benzoic acid as a cocatalyst (20 mol%) under the best reaction conditions. The product 7k derived from the reaction of 5b with 6a was obtained in good chemical yield, moderate diastereoselectivity and enantioselectivity (84%, anti: syn/64:36 dr, anti: 60% ee, syn: 57% ee). The product 71 derived from the reaction of 5b with 6b in good chemical yields and stereoselectivities (82%, anti: syn/ 90:10 dr, anti: 74% ee, syn: 63% ee). Whereas, the reaction of 7 membered 5c with 6a did not proceed well giving the aldol product 7m in low chemical yield and diastereoselectivity with poor enantioselectivity (32%, anti: syn/50: 50 dr, anti: 14% ee, syn: 4% ee). Moreover, the use of substituted cyclohexanone 5d also afforded the product 7n with moderate chemical yield and stereoslectivities (62%, anti: syn/53:47 dr, anti: 58% ee, syn: RSC Advances Paper

49% ee). In addition, the reaction using pyranone **5e** as heterocyclic ketone was also examined and the aldol product **7o** was afforded in good chemical yield and diastereoslectivity with moderate enantioselectivity (82%, *anti*: *syn*/60: 40 dr, *anti*: 43% ee, *syn*: 42% ee). The reaction of acetophenone **5f** with **6a** afforded the aldol product **7p** in good enantioselectivity but moderate yields (60%, 94% ee). The reaction of acyclic ketones **5g,h** with **6a**, respectively, gave the aldol products **7q,r** as racemate with low chemical yields. The aldol reaction of cyclohexanone with aliphatic aldehydes like **6k** and **5g** as acceptors, was also tried but the desired product was not observed.

2.5. Reaction mechanism

Considering both the good enantioselectivity (94% ee) of the chiral aldol product [2S,3R]-7a that was obtained in the reaction of 5a with 6a and the results of its calculation studies (Fig. 1–3), an enantioselective reaction course is proposed as follows (Scheme 5).

For the estimation of enantioselective reaction course, initially the conformational analysis using the scan of total energies for the enamine intermediate I was carried out and the result indicated the conformation of enamine intermediate I-1, with an intramolecular hydrogen bonding between hydrogen



Scheme 5 Plausible reaction course of the reaction.

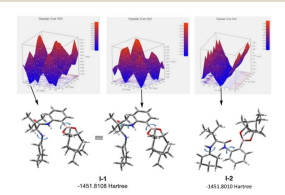


Fig. 1 Scan of total energies (up) and DFT optimized structures (down, at the B3LYP/6-31G(d) level of theory) of the enamine intermediate I generated by varying the torsion angles (the dihedral scans showed with u-shaped arrows: I-1).

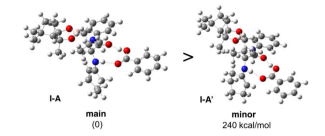


Fig. 2 The scan of total energies for I-A and I-A'.

atom of amide amino group and oxygen atom adjacent to boron atom, suggesting I-1 is better than I-2, as I-2 does not have similar intramolecular hydrogen bonding (Fig. 1). Furthermore, when benzoic acid as a co-catalyst is coordinated to more stable I-1 by hydrogen bonding, it is suggested that I-A, in which I-1 and benzoic acid are hydrogen bonded at two points to each other, is more stable than I-A' (Fig. 2). In addition, it is also indicated that in I-1, the olefin of the enamine was anterior, but in I-A, its olefin has a posterior conformation (Fig. 3). Moreover, to examine the regioselectivity of the reaction between I-1 and 6a, the calculation of the energies (Fig. 1 and 3) and coefficients of their frontier orbitals (Fig. 4) were conducted. The energy levels of the orbitals calculation clearly showed the interaction between the LUMO of 6a and the HOMO of I-1, and their orbital phase clearly demonstrated a matching in favor of overlapping to afford the observed configuration of major aldol product 7a (Fig. 4).

Based on the above calculation results, the reaction might pass through enamine intermediate I-1 and complex I-A (I-1-

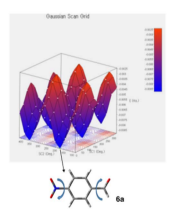


Fig. 3 Scan of total energies (up) and DFT optimized structures (down, at the B3LYP/6-31G(d) level of theory) of **6a** generated by varying the torsion angles (the dihedral scans showed with u-shaped arrows).

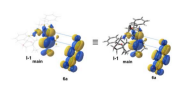


Fig. 4 The frontier orbital between I-1 and 6a.

benzoic acid) and subsequently, it was indicated that the reaction of I-A with 6a might progress through proposed four different transition states Ts-1-4 as shown in Scheme 5. However, among the proposed Ts-1-4, the reaction might proceed through Ts-2, based on the scan of total energy of I-A (Fig. 2) and the frontier orbital analysis (Fig. 3), having smaller steric interaction both between I-A and aldehyde 6a than those of Ts-1, 3, 4.

In contrary to the expectations, the calculation results suggest that, the boron atom on the catalyst does not coordinate the substrate **6a** as a Lewis acid moiety. However, the boron atom has three covalent bonds in the plane, and its configuration may contribute to the formation of preferable transition states in affording aldol product **7a** with a high stereoselectivity. Furthermore, it is also suggested that boron atom might coordinate with water solvent, forming an efficient conformation of the transition state, affording high enantioselectivity, although it is not clear why the solvent system is effective.

3. Conclusion

Paper

We have developed a new boron fused primary amino amides and their functionality as organocatalysts were examined in the asymmetric cross aldol reactions of various ketones 5a-h with different aromatic aldehydes 6a-k to afford the desired chiral aldol products 7a-r. The developed catalysts showed catalytic activity in this reaction. Particularly, catalyst 4b showed good catalytic activity to afford the various chiral aldol products in satisfactory chemical yields and stereoselectivities (up to 94%, anti:syn/68:32 dr, up to 94% ee) and interestingly, the effective solvent of the reaction was the mixture of sea H_2O and tap H_2O .

4. Experimental

4.1. General information

All reagents and dry solvents were purchased from commercial vendors and used directly without further purification. All reactions were placed in dried sample vials inserted with magnetic beads. ¹H and ¹³C NMR spectroscopic data were recorded using a JEOL JNM-ECA500 instrument with tetramethylsilane as the internal standard. Thin-layer chromatography (TLC) was performed on Merck silica gel 60 F254 plates and the analytes were identified under UV light. Flash column chromatography was performed using silica gel pore size 60 N (40-100 μm). Infrared (IR) spectra were measured with a JASCO FT/ IR-4100 spectrophotometer. HPLC data were collected using the TOSOH instrument equipped with (UV-8020, DP-8020, and SD-8022) detectors using Daicel Chiralpak AD-H, AS-H, OD-H columns. Optical rotations were measured with a JASCO-DIP-370 digital polarimeter. MS were taken on a JEOL-JMS-700V spectrometers.

4.2. General procedure for catalytic asymmetric aldol reaction of various ketones 5 with aromatic aldehydes 6

Catalyst **4b** (10 mol%) and co-catalyst (20 mol%) were added to a solution of ketones (0.4 mmol) and the respective aldehydes

(0.1 mmol) under the sea $\rm H_2O$ -tap $\rm H_2O$ (1:1) as solvent reaction condition. The reaction is allowed to stir at room temperature for appropriate time until the reaction completion. After the reaction completion, the solvent was removed under reduced pressure and the residue was purified by flash chromatography on silica gel (n-hexane/CH₃CO₂Et) to give the corresponding aldol products. The compounds are the known compounds and the structures were identified by spectral data which were in good agreement with those reported. The enantiomeric excess (ee) was determined using high pressure liquid chromatography (HPLC) principle by Daicel Chiralpak AD-H, AS-H, OD-H columns. $^{11a-q}$

Conflicts of interest

There are no conflicts to declare.

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